

UCAM & UNAH

Peru Project

Fieldwork & Laboratory

Water Manual



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Introduction:

Organising the field data collection

***The tasks -**

- a. Water samples:** the number of samples to be collected depends on the site location.
- b. Sediment samples (suspended load, bed & bank):** two samples per site – bed and bank.
- c. Water quality tests:** pH, conductivity, water temperature, total dissolved solids, dissolved oxygen & oxygen reduction potential.
- d. River channel data:** width, depth & velocity.
- e. Atmospheric data:** air temperature & air pressure.

***Division of tasks –**

The data could be collected by a single team or two teams. If the latter, then the tasks could be divided between the two teams as follows -

- Tasks **a**, **c** and **e** could be undertaken by one team
- Tasks **b** and **d** could be carried out by the other team.

Section 1: Equipment calibration

Equipment calibration is the most important step to ensure the accurate measurement of parameters. Check each item of equipment for correct functioning and calibrate according to the following procedures **before** starting a field visit.

NB. ensure that all batteries are installed and charged, and that spares are available.

Safety: refer to the Risk assessments in the Appendices for safe handling of equipment.

1.1. Hanna Multiparameter (HI98494)

Measuring: pH, Electrical conductivity (EC), Dissolved oxygen (DO), Total Dissolved Solids (TDS), Oxygen Reduction Potential (ORP), Atmospheric pressure & Water temperature.

Equipment & Materials:

Hanna Multiparameter (HI98494) & Accessories; Allen key (to remove probes); pH probe spare; spare DO probe; spare EC probe; spare DO Smart caps; Conductivity calibration solution (HI70031); Quick calibration solution; pH 4.01 buffers (HI70004P); pH 7.01 buffers (HI70007P); pH 10.01 buffers (HI70010P); Batteries (AAA); Distilled water; Electrode storage solution (HI70300); Sodium sulfite (Na_2SO_3), beaker (250+ml)



Check and take action:

***Allen key:** available to remove/tighten probes.

***Battery life:** automatically checked. Check & recharge every few months.

***Cleaning & Calibration:** see below.

Calibration:

i. Unscrew the storage container and **dispose** of any liquid in it.

ii. **Remove** the caps on any of the probes and **rinse** the probes with distilled water.

Note: after prolonged non-use (several months) the pH/ORP probe will need changing.

Note: after prolonged non-use (ie.2+ months) the DO Smart cap will need changing – follow instructions that come with the new ‘DO Smart cap’.

iii. **Use** the brush provided to **remove** any deposits/sediments on the probes/sensors that have accumulated during storage.

Note: Salt deposits may accumulate on the pH/ORP sensors after long periods of storage. This should be removed completely using the brush with distilled water. It may be necessary to leave the probe standing in a sodium sulphite solution* (200ml) for an hour, if the deposits remain.

*Solution: mix sodium sulphite (1gr) per 100ml of distilled water and leave for 10 mins to dissolve.

iv. **Turn on** the multiparameter by pressing the red/white button.

v. **Confirm** that the multiparameter is reading all probes/sensors one-by-one by pressing the button with the white dot (top left).

Use the arrows to **scroll down** and check that the probe/sensor is registering each reading.

vi. Press '**Escape**', then **Select 'Menu'** and scroll down to '**Calibration**'.

vii. **OPTION 1: Select 'Quick calibration'**

Note: only select this option if the equipment has been used recently, or just checking. If it has not been used for many weeks or months, then a full calibration is required.

***Fill** the storage container/wide test tube: wide enough to accommodate all the probes, so that it is one third full with the '*Quick calibration solution*' and place the probes in the container.

***Wait** until it is **confirmed** on the screen that each probe/sensor is calibrated (ready).

***Remove** the probes/sensors from the container and hold them in the air until it is again **confirmed** on the screen that each probe/sensor is calibrated (ready).

NB. This is especially important for the calibration of the dissolved oxygen probe - hold the probe very close (almost touching) the water surface to calibrate for humid air.

***Select 'Tick'** (top left – white dot button) when done. The multiparameter is ready for use.

***NB.** During more or less **daily use**, **fill** the storage container with tap water and refit to the device so that the probes are kept moist.

If it is **not** going to be used for **many weeks**, then follow the **Storage procedures** in **Section 5**.

viii. **OPTION 2: Select 'Single param. Calibration'**

***Select** the parameter of interest: pH, ORP, Conductivity or %DO and follow the instructions using the correct buffer fluids.

NB. '**Factory reset**' should be **selected** prior to selecting each parameter.

-**pH:** select "**Calibrate pH**" and then '**Factory reset**' - **calibrate** following the instructions on the screen using up to three buffers (4.01, 7.01, 10.01). One third **fill** the storage container or another suitable container with the required buffer so that it would cover the probe and **place** the probe in the container. Follow the 'instructions' on the screen until it displays '**ready**'.

Repeat with the next buffer until all three buffers are calibrated and the screen displays '**ok**'.

-**ORP:** separate calibration is **not** required for a new ORP sensor because it is on the pH probe and they can be calibrated together, but doing so does establish a baseline that can be used as a comparison for future validations. Calibration compensates for changes due to contamination of the platinum surface and drift in the reference electrode. (See p34 in Hanna Manual).

NB. A relative mV calibration can also be made to remove the voltage attributable to the Ag/AgCl reference electrode (to display the ORP versus a SHE (standard hydrogen electrode)). This is really an arithmetic correction and is correct only at the standard temperature. For example, test solution HI7022 reads 470 mV at 25 °C versus the Ag/AgCl reference. The ORP mV versus a SHE would be 675 mV (add 205 mV to the observed value).

-**EC:** select "**Calibrate Conductivity**" and then '**Factory reset**' - **calibrate** following the instructions on the screen using the 1413µ solution. One third **fill** the storage container or another suitable container with the required buffer so that it would cover the probe and **place** the probe in the container. Follow the 'instructions' on the screen until it displays '**ok**'.

-%DO: it is best to calibrate on arrival at the general location of deployment (ie.in Huanta). Errors in measurement may result if altitude and barometric pressure differ greatly between the calibration location and measurement sites. (See Manual page 36).

-to calibrate to 0%: place the DO probe in the container that has been two thirds filled with **Zero oxygen solution**. The DO probe takes time to **stabilise** but should do so at or close to '0%'. **Remove** the DO probe from the liquid and **shake** gently to remove excess fluid. **Hold** just above the solution and it should stabilise at approx. 70%. **Reimmerse** the probes in the solution to confirm the DO % as 0%.

-to calibrate to 100%: place a little **distilled water** in the container. **Place** the probe in the container but do not tighten and wait 10+ minutes before **pressing** 'Start' to begin the calibration.

***To replace the DO Smart cap** (See Hanna Manual page 16)

NB.with a new probe, Hanna recommend immersion in the Zero oxygen solution for up to 8 hours!

NB.ensure that the Zero oxygen solution is mixed correctly: it may be supplied in two bottles that need to be mixed to create the solution.

NB. sodium sulfite solution is an alternative to using a zero oxygen solution. Mix 1gr of sodium sulfite with 100ml of distilled water. Follow the instructions above for 0% and 100% calibration.

1.2.Mettler Toledo (FiveGo)

Measuring: Dissolved oxygen (DO).

Equipment & Materials: Mettler Toledo (FiveGo), distilled water.

Check & Take action: check battery status and that the cap contains distilled water when not in use.

NB. Confirm the air pressure and calibrate at each site.

i.**Turn** on the equipment.

ii.**Air pressure setting:** this is important when measuring dissolved oxygen and it will **vary between each site**.

-Press 'Setup' until the air pressure value in mbars appears. **Adjust** the value by using the arrow tabs to the air pressure value given by the multiparameter.

-Press 'Read' to confirm the value entered.

iii.**Calibration** - a one point calibration is suited to water with an oxygen concentration over 1 ppm.

-Hold the sensor in the air and **press 'Cal'** and when the screen displays '**100%**' press '**Read**'.

iv. **Place** the sensor in the channel and **press 'Read'**. When the letter '**A**' stops blinking and is stable, **record** the value. **NB.**this may take some time.

v.**Measuring DO** -

-Use the 'Mode' button to switch between '**%**' and '**ppm**'.



Section 2: WATER & SEDIMENT SAMPLING

Safety: Always wear gloves and appropriate PPEs when taking samples. Refer to the labels on bottles/tubes for chemical preservatives in them and read the risk assessments.

Important: The aim of collecting water and sediment samples is to obtain samples that are true representations of the water or sediment at the point of collection. That means preserving the sample according to the guidelines given below and avoiding any possible contamination.

2.1. Water sampling

-Collect water samples at each site in new or pre-cleaned HDPE bottles (1000ml).

-Rinse bottles for reuse with hydrochloric acid (1%) solution, if available, and/or distilled water until pH7. Use pH papers to check that the bottles are pH neutral (pH 7.0).

-Collect the water samples using the extendable bucket from the middle of the channel just below the surface, if possible.

-Rinse the bucket and HDPE bottles with water **3 times** prior to collecting each sample.

-Prepare subsamples as follows:

- For **Anion & cation tests** and **general** tests, water samples need to be filtered. **Collect** water into HDPE (1 litre) bottles and then **filter** the quantity required into the PP sample tubes or bottles provided by SGS.
- For **Metals** the sample should be filtered for comparison to *regulatory levels* with 0.45µm filters. For *hydrochemical analysis* the water should be filtered with 0.22µm filters. For Metals, **nitric acid** should be **added** to the sample as soon as practical to reduce it to pH2 - see below.
- For **COD, biological, pesticide, total nitrogen** and **total phosphate** tests, water samples should **not** be filtered. **Collect** the samples directly into the PP sample bottles provided. **NB.Pesticide** sample bottles should be filled to the brim of the bottle. **NB.COD** sample bottles should have sulphuric acid added to reduce the sample to pH2.
- For **Alkalinity** tests in the lab., unfiltered water in the HDPE bottles should be used.

Notes:

-**Nitric acid** needs to be used to stabilise the samples for **metal** tests. The acid should be **ultrapure** (<99.99% trace metal basis) or distilled in the lab (double distilled is better but single distilled will be OK). If concentrated nitric is used, the molarity is about 15M and only 2-4 drops are required per 60ml of sample. The sample should have a **pH** as close to **2** as possible. **Check** with pH strips.

For practical purposes and safety, add the nitric acid to the test tubes in the lab.

-If nitric acid is not available, the sample must be **filtered in situ** directly in to the test tubes. Samples should be transferred to the analytical lab within two weeks, where the metal and cation samples should then be acidified to ensure all adsorbed or precipitated materials return to solution.

NB. If **Anion and cation** samples are to be analysed **together** then do **NOT** acidify the sample. This will nullify the anion analysis.

-Bottles supplied for pesticide tests may contain *sodium thiosulphate* which can be an **irritant** (See **Appendix: Risk assessment for sampling preservation**).

-**Always use** the correct bottle/tube for samples as they are specific for the component being tested. Plastic tubes cannot be replaced with glass ones and vice versa. Chemical preservatives should not be allowed to cross-contaminate the samples.

-**Do not use** bottles/tubes that are not supplied for this field visit. They may contain expired chemicals or not have been cleaned for the purpose, causing contamination of the samples.

-**Do not rinse** previously unused sample test tubes/small bottles with distilled water prior to introducing the sample.

Preparing filtered samples:

Filtration is used to remove turbidity due to any organic and inorganic particles / sediments and biological materials to preserve the sample from degradation. Filtration collects and preserves the filtrate on a filter paper.

A. Using a filter holder

Equipment & Materials: Swinex filter holder; gaskets; 0.22µm (*hydrochemical analysis*) or 0.45µm (*regulatory levels*) filter membranes; distilled water; tweezers; petri dishes;

Equipment set-up:

-**Separate** the two sections of the filter holder and **rinse** inside and out with distilled water.

-**Separate** a white filter paper (0.22µm or 0.45µm) from the blue protective disks/covers, without damaging the filter, and **place** inside the filter unit using tweezers.

-if there is wording or a grid printed on the filter, it should be facing upwards.

-**Ensure** that the rubber 'O' gaskets (not shown in the images below) in both halves of the filter unit are in place and **screw** the two sections together.

-**Attach** the filter holder to the tubing for pump or to the syringe following Sections B and C.

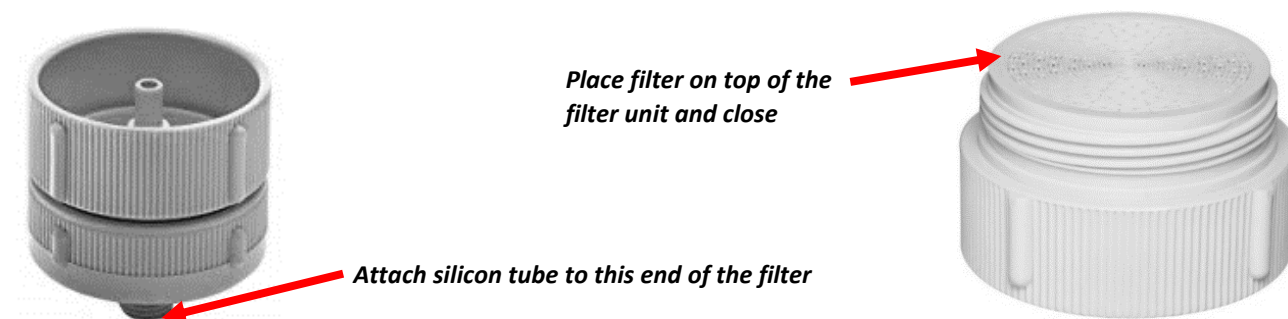


Figure: (left) Swinex filter holder with two sections screwed. The inlet tubing/syringe is connected to the bottom (See photo above); **(right)** Swinex filter holder with the filter holding section opened.

After filtering samples from one site:

-**Weigh** the **empty bottle** before pumping the water via the filter into it. **Weigh** the bottle again at the end of the filtering process. **Record** both **weights** on the field data sheet and **calculate** the quantity of water that has passed through the filter.

-**Open** the filter unit carefully and **remove** the filter paper using tweezers.

-**Place** the filter paper in a petri dish (keep **all** filter papers used at each site, if more than one used). The filter papers trap suspended load which can also be analysed.

-Clearly **write** on the petri dish the site no., name, date and the amount of water that has passed through the filter / filters (eg.813ml) and record on the data sheet.

NB. If more than one filter paper is used due to the large amount of sediment present, follow this procedure for all filter papers used. Once dry, several filter papers can be stored back to back in one petri dish.

-**Use** distilled water to **clean** the filter unit before and after filtering water from **each site**.

-**Leave** the filter paper / papers to **dry naturally overnight** (lid partially covering) and **seal** the petri dish / dishes with parafilm the **next day**.

B. Using a peristaltic pump

Equipment & Materials: Geotech peristaltic pump; AC/DC combination power cable; rechargeable battery; DC power cable with car battery connectors; silicone tubing; distilled water; HDPE bottle (1000ml); Swinex filter holder with a new filter paper inserted (See section A above); tubes/bottles for collecting filtered samples; HDPE bottles (1+ bottle).



***Equipment set-up -**

***Check** the pump 'on/off' button is in the 'off' position. **Connect** the power cables to the 12VDC supply at the back of the pump;

***Lift** the lever in the pump head and **move** fully to the left to open the pump head;

***Place** the tubing through the pump head;

***Move** the lever to the right to lock and **close** the pump head with the tube within it;

***First** pass distilled water through the tubing to clean it and ensure it is working smoothly.

Then, pass a small amount of sample water through the tubing before attaching the filter unit.

***Attach** the Swinex filter holder firmly, containing a **new filter paper**, to the end of the tubing.

***Filtering the sample -**

- Collect** a water sample in an HDPE bottle (1000ml) in the field.
- Place** one end of the silicon tubing in to the sample collection bottle.
- Place** the other end of the silicon tubing in the empty HDPE bottle.
- Set** the pump direction to 'forward' using the 'reverse/forward' button.
- Turn on** the pump and **set** the speed to 3-4 (marked on the dial).
- Towards the end of the filtering, **ensure** that all sediments in the water sample are **extracted** through the filter by gently shaking the sample bottle.
- Once the new sample collection bottle is filled, **reweigh** the sample collection bottle and **calculate** the weight / volume of filtered water in the bottle.
- Label** the new sample collection bottle containing the filtered water with the **site no.** and **date.**
- Use** the bottle of filtered water to fill each of the sample bottles (ie.metals) with the required amount of filtered water.

***Filter becomes clogged -**

- If the filter becomes **clogged** with sediment, water will no longer pass through it.
- Open** up the filter unit, **remove** the filter paper and **replace** with a new one.
- Do **NOT clean** the filter unit.
- Store** (See below) all the filter papers (a river with a lot of suspended sediment may require 6+ filter papers to filter 1000ml).
- After collecting sufficient filtered water for the site, **turn off** the pump.

***Storing the filter papers -**

- Open** the Swinex filter.
- Remove** the filter paper with tweezers and **place** in a petri dish labelled with site no., name, date and the **total weighed amount** of water that has passed through the filter (eg.813ml).
- Leave** the filter paper to **dry naturally overnight** (lid partially covering) and **seal** the petri dish with parafilm **the next day.**

***Cleaning the filter unit –**

- Remove** the filter unit from the tubing and **clean** fully both sections of the unit with distilled water (before filtering the sample from another site).

***Cleaning the silicon tubing -**

- Place** one end of the tubing in to a bottle of distilled water.
- Turn on** the pump while keeping a waste container below the end of the tubing to collect the water that is flowing through the tubing and cleaning it.
- The tubing can be reused with samples from different sites as long as this procedure is followed and the tubing shows no mechanical damage.

Note: keep the tubing clean and dust free during transport between sites, if using in the field.

C. Using a syringe to filter:

Equipment & Materials: 50ml syringe (1 per site); Swinex filter holder with a new membrane inserted (See Section A above); tubes/bottles for collecting filtered samples; HDPE bottles (1+)

- Fill** a new syringe with the required amount of water from a HDPE bottle;
- Attach** the syringe to the top inlet of the Swinex filter;
- Place** a clean PP tube of HDPE bottle for collecting the filtered sample and compress the syringe until the required amount of water has been filtered;
- Once the water sample has been added to the Sample collection tube, the cap must be fully tightened and then secured with parafilm;

Note: Use a new syringe for each site since proper cleaning of the syringe may be difficult.

Water samples collection for testing for 'heavy metals'

For the water samples destined to be tested for 'heavy metals', the water sample must contain 0.5ml or 3+? drops of nitric acid (70%, 99.99% trace metal basis) as appropriate to reduce the water sample to a **pH of 2**.

Note: Past experience indicates that changes in atmospheric pressure at altitude can cause **nitric acid** to **escape** from the test tubes, if the test tubes have been pre-filled.

In this case, the following procedure should be followed –

- a tightly closed bottle of nitric acid should be carried, regularly monitored and **carefully stored** during field visits.
- one person should be designated and responsible for handling the nitric acid bottle at all times.
- nitric acid should only be added to test tubes under controlled conditions, in line with the all-risk assessment procedures.
- once the water sample and nitric acid have both been added to the test tube, the cap should be tightly fitted and then secured with *parafilm*.

2.2. Sediment sampling

A. Sediment sample collection

Equipment and materials: plastic sample containers or plastic bags (small – 2-3 per site); spoon/spatula/trowel; marker pen; gloves;

Sample collection:

- Collect** sediment samples at all sites if possible, from the river **bed** (permanently under water) and **banks** (rarely under water);
- The sample should consist of **finer** material: silt/sand. If possible, take the sample from **just below the surface** of the river bed or bank. Fill the bag to a depth of about 5cms (1kgs);
- River bed:** **scoop** a sample from as far out in to the channel as is practical (safe);
- River bank:** **scoop** a sample from a place **higher** up the bank or lakeside that is clearly affected occasionally by the river when it is in flood;

- Leave** the container/bag to 'rest' and then **drain** out most of the water prior to sealing;
- In the lab. a subsample of 50+grs can be extracted and the rest archived for future use.
- Seal** the container/bag and clearly **label** the container with the site no., site name, date and time of collection.

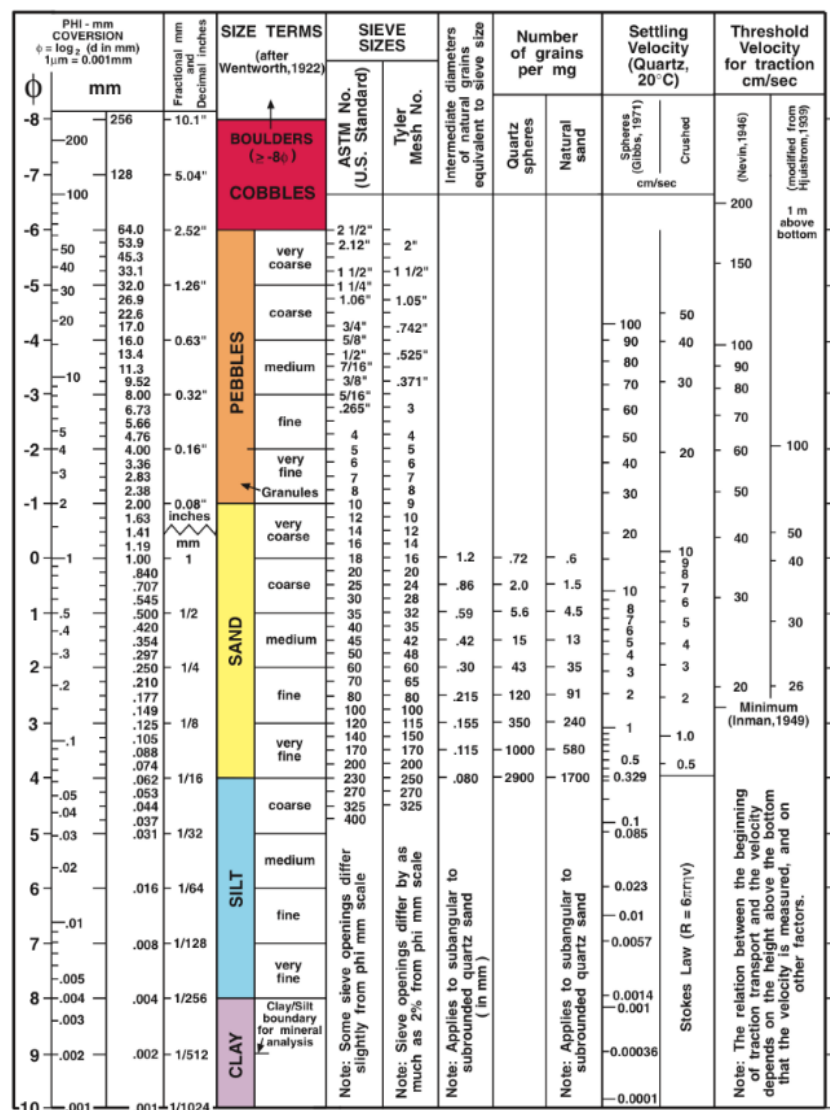
B. Sediment size recording

Sediment size data is recorded by observation with reference to the **Wentworth sediment scale** chart – please see below.

An estimation should be made of the approximate percentage of the main sediment types (qualitative not quantitative) occupying the river channel. The terminology as stated in the Wentworth sediment scale chart should be employed.

This data can be double checked subsequently by referring to photos taken of the channel.

Material	Size
Sand	<2mm
Gravel	2mm – 15mm
Pebbles	15 – 60mm
Cobbles	60 – 250mm
Boulders	250mm+



Section 3: FIELD MEASUREMENTS

3. 1. Physico-chemical measurements

Parameter such as pH, Electrical conductivity, Temperature, Dissolved oxygen and ORP in water samples depend on many environmental factors including air pressure, temperature and bacterial activity. These parameters are therefore highly susceptible to rapid change after collection. Measurements should be taken on-site to ensure accurate values for these parameters – they are displayed across three screens on the multi-parameter.

All meters should be calibrated according to **Section 1** before measurements are taken.

1. Hanna Multiparameter (HI98494)

Measuring: pH, Electrical conductivity (EC), Total dissolved solids (TDS), Dissolved oxygen (DO) & Water temperature.

Equipment & Materials: Hanna Multiparameter (HI98494) (calibrated); distilled water; tissues;

Equipment set-up:

-**Clean** the electrodes with distilled water, ensuring that the water enters the gap at the end of the sensor, to remove any salts from the saline solution it is stored in.

-The probes (electrodes) can be directly immersed in the river/lake for measurements.

A. pH measurement:

***Measurement unit:** pH

-**Place** the end of the sensor in the water, ensuring that it is completely immersed in the water.

-**Select** the 'pH' mode.

-**Wait** until the 'unstable' display disappears and then **read off** the result.

B. Electrical conductivity (EC) measurement:

***Measurement unit:** Low range ($\mu\text{S}/\text{cm}$), High range (mS/cm)

-While the end of the sensor is in the water [See above], **select** the 'conductivity' mode.

-**Wait** until the 'unstable' display disappears and then **read off** the result.

C. Total dissolved solids (TDS) measurement:

***Measurement unit:** High (ppt) / Low (ppm)

-Keeping the sensor in the water [See above], **select** the 'TDS' mode.

-**Wait** until the 'unstable' display disappears and then **read off** the result.

D. Dissolved Oxygen measurement:

***Measurement unit:** % and mg/l of DO.

-Keeping the sensor in the water [See above], **select** the '**DO**' mode.

-**Wait** until the 'unstable' display disappears and then **read off** the result. This may take some time.

E. ORP measurements:

***Measurement unit:** mV

-Keeping the sensor in the water [See above], **select** the '**ORP**' mode.

-**Wait** until the 'unstable' display disappears – this may take about 5 minutes - and then **read off** the result.

F. Water temperature measurement:

***Measurement unit:** °C (degrees centigrade)

-Keeping the sensor in the water [See above], **select** the '**Water temperature**' mode.

-**Wait** until the 'unstable' display disappears and then **read off** the result.

G. Atmospheric pressure (Atm Pressure) measurement:

***Measurement unit:** mbar

-The atmospheric pressure is displayed when the meter is turned on. **Read off** the value.

After measurements:

-**Clean** probes with distilled water after use.

-**Replace** the plastic cap on the **pH sensor** immediately to protect it.

-**Storage:** when the use of the multiparameter is complete, **place** a little storage solution (1cm) in the storage container to keep all the sensors moist and screw the storage container on to the multiparameter.

2. Measuring Alkalinity

A. Alkalinity Test strips

Equipment: Test strips (ie.Lamotte)



-**use** like pH strips and dip in to the sample. The test strips are designed to be read off immediately (2 secs) after they have been dipped in the sample.

-**record** which of the options (ie. 0, 40, 80, 120, 180, mg/L (ppm)) is indicated on the scale.

-if the colour is trending towards the next point on the scale, this suggests that the true result might be between the two points on the scale, **record** the lower reading with a '+' sign.

B. Alkalinity tester

Equipment: Hanna (HI-775 Freshwater Alkalinity Colorimeter - Checker®HC), Reagent kit, Batteries (AAA x 2)

-‘Zero’ the Checker®HC by **placing** the empty vial inside the checker and **pressing** the button (optional).

-**remove** the vial, **add** the water sample up to the ‘white’ line on the vial and **press** the button.

-**clean** and **dry** the vial before placing it in to the tester.

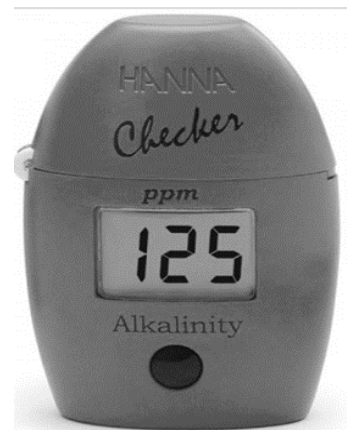
-when prompted to do so, **remove** the vial and **add** 10ml of the **reagent** using the pipette provided.

-**clean** and **dry** the vial before placing it in to the tester.

-**place** the vial back into the Checker®HC, **press** the button and read off the result.

-after each use, **clean** the vial thoroughly with distilled water. **NEVER** clean inside the tester.

NB. Check after each visit that there is sufficient reagent.



3.2. River discharge rate measurements

Safety: Avoid stepping into the river whenever possible. Measurements should be taken while in the river water only if the river is shallow, the bed is sturdy and the flow rate is slow.

Theory: The water discharge rate is calculated using the velocity (v_1 - v_5) and depth (d_1 - d_5) measured at equal intervals along a selected cross-section (See figure below) of the river.

For this, first select a suitable cross-section of the river. Take a width measurement, calculate the intervals for depth and velocity measurements, and measure the velocity and depth at these intervals (See figure below).

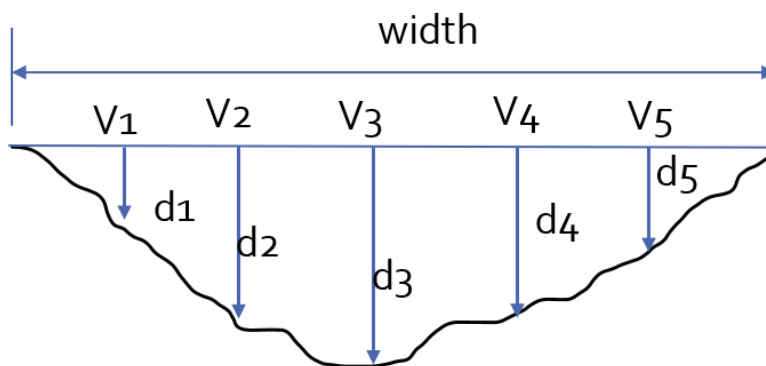


Figure: A cross-section of river showing the points where the measurements need to be taken

1. Width measurement

These measurements can be made using a tape measure or a telescope laser distance meter.

***Measurement: m (metres)**

A. Using a tape measure:

-Place the tape measure at a **90° angle** to the river bank.

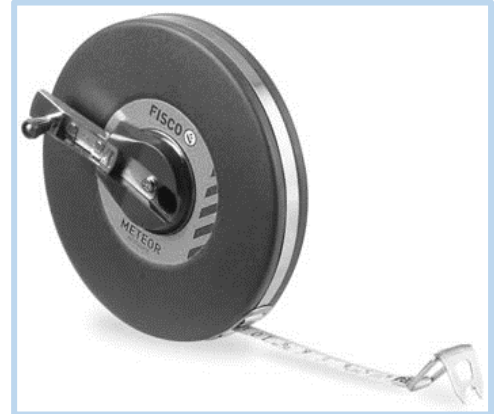
-Measure from the point on one side of the river where the **water touches the bank** to where it touches the bank on the other side, **across the surface** of the river.

-If the bank is **muddy**, a decision will need to be made as to where the surface of the water is touching the firm bank. Too shallow water that is not flowing can be neglected from the width measurement. Note this in the data recording sheet.

-If the bank has an **overhang**, the tape measure must be placed under the overhang to where the surface of the water is touching the firm bank.

-Ensure that the tape is pulled tight **just above the surface** and is not dragged under water.

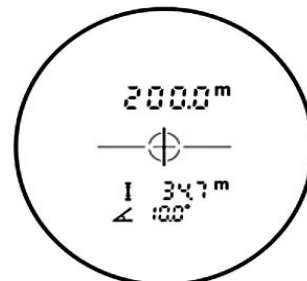
-If the width of the river is composed of **more than one channel** (braided), then the width of all channels needs to be measured to obtain the total width. In that case, measure the flow rate and depth of each channel as well to calculate the total discharge of the river at the site.



B. Using Laser Works monocular telescope laser measurer:

This method is useful when it is difficult to reach the opposite bank of the river.

Theory:



- The meter measures both height and distance if the laser is projected at an angle. For river measurements, **keep the laser angle to zero.**

Method:

- Turn on the measurer.
- Stand on the edge of the river bank, in-line with where the **water touches the bank**.

Note: You should crouch down to obtain a proper alignment just above the surface of the river.

- Hold** the measurer **very steady** and press the 'mode' button until a 'diamond' appears in the centre of the screen.

- Aim** the measurer at a **90° angle to the bank** (the display should show **0°** laser angle with the ground) at the opposite bank where the water touches the bank and position the 'diamond' in the centre of the screen on a rock just above the water level on the opposite bank.

- Press** the 'on/off' button once and the **width**, in metres, is displayed on the screen.

- Repeat** 2-3 times to confirm the result.

*See also the guidance in **Section A** above.



2. Depth measurements

*Measurement: m (metres)

One depth measurement will not give a true measure of the depth across the river channel. Consequently, it is best to take 5 measurements (3 if width less than 1 metre) at equal intervals along the selected cross-section to calculate the discharge rate as described in Section 3.2.

- Divide the width measurement by **six** to obtain **five** locations at which to take **five depth measurements** at equal intervals across the river (Example: width = 6.0 m; $6.0/6 = \text{every } 1.0 \text{ m}$).

A. Using a metre ruler

- Use a **metre rule** to measure the depth of the water. Extension pole or the flow meter pole may also be useful in conjunction with a tape measure.
- Do **not** push the ruler into the river load/bed material.
- Hold the metre rule **vertically** and read off the depth.

B. Using a Geopacks or Valeport Flowmeter

- Use a flowmeter pole to measure the depth of the water: the Valeport flowmeter has a scale on the pole.
- Do **not** push the flowmeter into the river load / bed material.
- Hold the flowmeter **vertically** and read off the depth.



3. Velocity measurements

***Measurement: m/s (metres per second)**

NB. the recordings should be taken at the same locations as the depth readings across the river.

A. Geopacks Flowmeter method: only suited to small channels.

-**Place** the flowmeter at **40% of the depth** down from the surface (60% of the depth up from the river bed).

-Make sure that there is no disturbance upstream (ie.someone in the river) and that the flowmeter **faces upstream**.

-The flowmeter must be held **vertically**, facing directly in to the current.

-**Hold** the flowmeter in position for **at least one minute** to note the average flow because at many sites the flow will vary over very short periods of time. **Record** the most stable velocity.

B. Valeport Flowmeter method: not suited to very large or very small channels.

-**Place** the flowmeter base on the **river bed**.

-Make sure that there is **no** disturbance upstream (ie.someone in the river) and that the flowmeter **faces upstream**.

-The flowmeter must be held **vertically**, facing directly in to the current.

-**Hold** the flowmeter in position to note the *average flow* because at many sites the flow will vary over very short periods of time. The Valeport averages the river flow over a specific time period set by the user (ie.15 seconds).

-Also, **record** the *standard deviation* given – a high figure indicates much variation in the flow while a low figure indicates little variation. The *standard deviation* should be significantly lower than the velocity.

4. Discharge measurements

a. Introduction

Calculating river discharge accurately in river/water studies is important. In water quality studies, discharge can be used to give accurate measurements of the absolute quantities of a given trace metal or sediments at a particular point at a given time in a river.

When using a flowmeter an important question to consider is **which section** of a river to use and **when** to take the measurements.

****Remember no river profile is uniform across its width and measurements always need to be taken from bank to bank.**

*It is best to use straight sections, away from riffles or pools. Try to ensure the riverbed is not very uneven with large boulders or in turbulent sections or section with eddies (to avoid backflow).

*If a discharge measurement is needed to correspond with water quality sampling, timing is not a consideration. If you want a baseflow measurement, it is best to take the measurements at least five days after rains cease. If storm discharge profiles are required, measure when the rains cease and thereafter, at five hourly intervals.

b. Methods & Procedures

***Field protocol:** always take river measurements with at least one other person present. If anyone asks what you are doing explain.

***Equipment needed:** flowmeter, tape measure, notebook, pens and camera.

***Clothing:** waders and rubber boots, depending on the river characteristics. A life jacket may be advisable, if a larger river is being measured.



Fig 1. Setting up flowmeter and tag line with measuring tape (Eustace Barnes)

i. In the field: at the river bank

Set the tape measure across the river ensure the tape measure is perpendicular to the river flow, is stretched tight and does not drag in the water. Fix with pegs to the bank. (See Fig.1).

ii. In the field: ready to start

You will be collecting data at regular intervals across the river. Those intervals are called stations. As a rule of thumb, stations should be about 1.0 metre apart on narrower rivers (up to 10 metres wide) and about 2 metres apart on wider rivers (10-20 metres wide). If there is a section where the river is deeper or the flow rate is higher, the intervals should be reduced. (See the suggested station intervals in Figure 2 below).

Feet	Meters	Number of stations
< 1.6	< 0.5	5 to 6
> 1.6 and < 3.3	> 0.5 and < 1	6 to 7
> 3.3 and < 9.8	> 1 and < 3	7 to 12
> 9.8 and < 16.4	> 3 and < 5	13 to 16
> 16.4	≥ 5	≥ 22

Fig 2. Best practice ideal 'station interval' across varying river widths

To start, face **UPRIVER**, just below the tape measure, so that the distances can be easily read. Place the flowmeter well in front of your feet so that readings are not affected by them. The river banks are to your left (LEW) and right (REW).

iii. Calculating river discharge –

Calculate an **average width** from *three* readings taken at the site (optional).

Calculate an **average depth** from all the readings taken at the site.

Calculate the **cross-sectional area (CSA): average width x average depth.**

Calculate the **average velocity** from all the velocity readings taken using one of the flowmeters.

Calculate the **discharge: cross-sectional area x average velocity.**

iv. Making an estimate of the river discharge -

Once you have gone through the above process, it is a good idea to see if the reading is about right. There are some easy field tricks that can be used to ensure your measurements are about right. However, this is no substitute for taking actual measurements.

NB. 1 cubic meter = 1,000 litres 0.5 cubic metres = 500 litres

So, what may appear a low reading, still represents a high channel flow.

*Worked examples -

For channels with the following widths: 2m, 5m, 10m, 15m and 30m.

- *with average depths of 0.2 and 0.8m*
- *with average velocities of 0.2, 0.25, 0.8 and 1.2m/s*

csa = cross sectional area; **ms** = metre squared; **mc** = metres cubed; **D** = discharge.

Channels 2m wide

- | | | |
|---|-------------|--|
| 1 (2m width, 0.2 depth, 0.25 velocity). | 0.4 ms csa. | D = 0.1mc/s (100litres per second.) |
| 2 (2m width, 0.8depth, 0.8 velocity). | 1.6 ms csa. | D = 1.28mc/s. (1280 litres per second) |

Channels 5m wide

3 (5m width, 0.2 depth, 0.25 velocity).	1.0 ms csa.	D = 0.25mc/s
4 (5m width, 0.8 depth, 0.8 velocity).	4.0 ms csa.	D = 3.2mc/s
5. (5m width, 0.8depth, 0.2 velocity).	4.0 ms csa.	D = 0.8mc/s
6. (5m width, 0.8depth, 1.2 velocity).	4.0 ms csa.	D = 4.8mc/s

Channels 10m wide

7.(10m width, 0.2 depth, 0.25 velocity).	2.0 ms csa.	D = 0.5mc/s
8. (10m width, 0.8depth, 0.8 velocity).	8.0 ms csa.	D = 6.4mc/s
9. (10m width, 0.8depth, 0.2 velocity).	8.0 ms csa.	D = 1.6mc/s

Channels 15m wide.

10.(15m width, 0.2 depth, 0.25 velocity).	3.0 ms csa.	D = 0.75mc/s
11.(15m width, 0.8depth, 0.8 velocity).	12.0 ms csa.	D = 9.6mc/s
12.(15m width, 0.8depth, 0.2 velocity).	12.0 ms csa	D = 2.4mc/s.

Channels 30m wide.

11. (30 m width, 0.2 depth, 0.25 velocity).	3.0 ms csa.	D = 1.5mc/s
12. (30 m width, 0.8depth, 0.8 velocity).	12.0 ms csa.	D = 19.2mc/s
13. (30m width, 0.8depth, 0.2 velocity).	12.0 ms csa	D = 4.8mc/s.

5. GPS co-ordinates of sites

GPS coordinates for each site are currently presented using decimal degrees on the UNAH UCAM website. However, many publications require authors to use degrees, minutes , seconds. GPS coordinates for each sample site can be obtained in two ways –

A.Using: GPS Finder - Garmin Etrex10

This is generally an inaccurate method because to obtain an accurate reading the GPS Garmin device requires contact with a minimum of 8 satellites. In much of Peru and especially in deeply incised valleys or remote locations, such as the Cañete valley, this is not possible. In these situations the GPS co-ordinates calculated for a site may be far away from the actual site. However, this method should be accurate where there is a wide flood-plain and contact with the satellites is possible.

Despite this, always record the GPS result. This can be corrected using Google Earth later (see Section B below) if accurate site data including photos, are also recorded.

1.Turn on the Garmin Etrex10 by pressing the '**Light**' button.

The screen opens with the cursor positioned over the '**Map**' tab.

2.Use the *toggle* to reposition the cursor over the '**Mark waypoint**' tab and **press** the *toggle* to select it.

3.Read off the '*Longitude*' and '*Latitude*' of the site (waypoint).

4.To obtain the **elevation**, it may be necessary to select the '**Mark waypoint**' tab a second time.

5.Use the *toggle* to move the cursor up to the '**Title**' at the top of the screen.

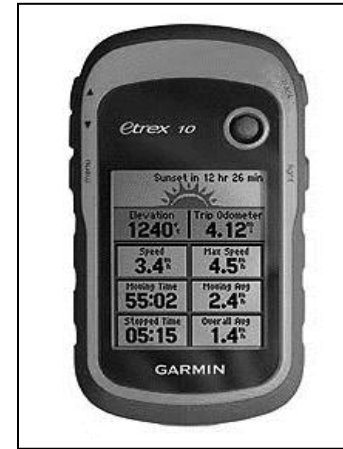
Press the *toggle* once and then – letter by letter, or number by number – enter the site name.

6.Move the cursor to the bottom of the screen when complete and select '**Done**'.

7.Move the cursor to '**Note**' and enter the site details (as above) as required.

8.Move the cursor to the bottom right of the screen and select '**Done**' – the data record will be stored.

9.To find stored waypoint data, select '**Waypoint Manager**' on the opening screen and scroll down to find the site/site data required.



B.Using: 'Google Earth Pro'

This allows the GPS co-ordinates for any site to be accurately determined. In the Google Earth Pro program, GPS data is shown automatically in the screen image information bar in the lower right hand corner of the screen, as follows: Imagery date, latitude, longitude, elevation and eye alt. (height).

The longitude and latitude information may be given in one of five ways:

- i.Decimal degrees;
- ii.Degrees, Minutes, Seconds;
- iii.Degrees, decimal Minutes;
- iv.Universal Transverse Mercator;
- v.Military Grid Reference System.

These can be selected on screen by going to 'Tools' on the task bar. In 'Tools' go to options and the five options are shown. Click on the option required.

1. Download and **open** up the '**Google Earth Pro**' app.

2. Search the required site location.

3. Position the cursor precisely on the site location.

4. Read off the **GPS co-ordinates** from the bottom of the screen.

5. Read off the **elevation** from the bottom of the screen.

Section 4: LABORATORY TESTS

Chemical tests for components that degrade quickly need to be carried out as soon as possible after the collection of samples. Alkalinity is best tested just after the collection of the sample. Nitrate tests must be done within 48 hours of the sample collection as long as proper storage conditions (acidified with 99.9% grade HNO_3 , at 4°C) are provided.

For practical reasons, it is recommended that both tests are carried out at the end of each day. Samples should be collected in bottles filled to the tip leaving no head space in them.

Safety: Always wear gloves, safety glasses and a Lab.coat during the tests. Refer to labels for chemical content and read the risk assessments for safe handling.

4.1. Titration Alkalinity Test

***Method:** Titration - to identify colour change at pH 4.5 (green to pink) in the sample.

***Measurement unit:** mg/l of calcium carbonate

***Equipment & Materials:** Digital Titrator; Titrator cartridges (1.6N & 0.16N); Phenolphthalein powder sachet; Bromcresol/Methyl powder sachet; 250ml flask; Distilled water; 100ml measuring cylinder; Magnetic stirrer; Scissors; Funnel;



***Sample collection:** collect 100ml of **unfiltered** water. **Maintain** in a cooler until ready to titrate but **raise** to room temperature for titration.

***Equipment set up:**

-**Insert** a cartridge in to the Titrator and twist 90 degrees to lock the cartridge in place.

Note: use a 1.6N cartridge where the water is more neutral (pH 6+) and a 0.16N cartridge where the water is very acidic (<pH 6)

-The cartridge can be used until it is empty.

Note: the cartridge contains sulphuric acid (H_2SO_4).

-**Remove** cartridge cap and insert detachable end tube.

Note: this must be changed if the cartridge is changed to a different concentration.

-**Press** in the sliding button and slide along shaft to expel any air in the detachable tube.

Note: at this point any drops of liquid must **not** be allowed to enter the water sample.

Note: there is a danger of acid shooting out of the detachable tube at this stage.

-For titration, first **zero** the reading using the large dial and then **turn** the button at the end of the titrator to dispense the drops of sulphuric acid (H_2SO_4).

-Once the test is complete, **use** the large dial to return the counter to zero.

***Equipment use:**

-**Use** a measuring cylinder to accurately measure 100ml of **filtered** water and transfer it into the 250ml flask.

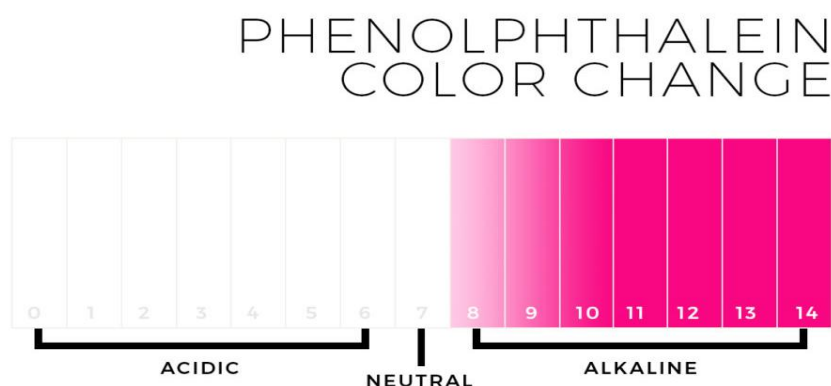
-Begin with the **1.6N** (high concentration) cartridge in the titrator unless water at the site is known to be especially acidic in which case use the 0.16N (low concentration).

***Test 1: Phenolphthalein test –**

-**Empty** the P.sachet in to the 100ml water sample and **shake gently** or place on the magnetic stirrer until completely **dissolved**;

-if the water goes even **slightly pink**, indicating a high level of alkalinity (pH 8.3+) – **start** titration - See instructions below (Point 3) - until the sample becomes transparent, then proceed to Test 2.

-If there is **no** change in colour, the water sample remains transparent, then proceed to Test 2.



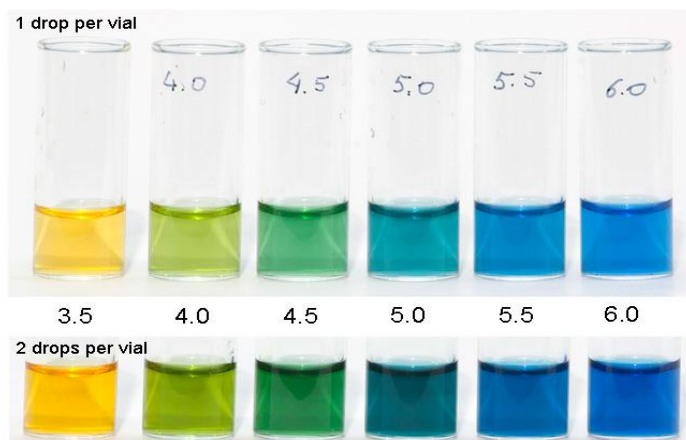
***Test 2: Bromcresol green / Methyl red test –**

- **Add** the Bromcresol sachet to the same water sample and **shake gently** or place on the magnetic stirrer until completely dissolved.

Note: the sample is most likely to turn **green**: Start titration. See instructions below (3).

Bromocresol Green pH Tester

pH Color Chart



*Test 3: Titration -

- **Ensure** that the titrator is set to **zero** (0000) and **no** air bubbles remain in the tube.
- **Place** the end of the titrator in to the flask but ensure that it does **not** enter the water.
- **Turn** the dial to dispense the sulphuric acid (H_2SO_4) slowly, one drop at a time.
- **Gently shake** the container or allow the magnetic stirrer to gently shake it as the sulphuric acid (H_2SO_4) is added.
- **Add** the sulphuric acid (H_2SO_4) until the sample changes colour (**1. Phenolphthalein test**: from pink to colourless; **2. Bromocresol green / Methyl red test**: green to light pink - until there is **no** longer a green/blue tint to the sample).

*Results:

- **Read off** and record the number displayed on the counter (mg/litre).

Note - if the number displayed on the counter is **<10** using the 1.6N cartridge when the sample turns light pink, then the cartridge must be changed to the 0.16N and Test 2 redone. The counter should **NOT** be set to zero but the score added to the original counter score.

After all tests are complete:

- **Dilute** the sample and dispose of it in to the **ground** - not in to water.
- Once all tests are complete, **press** and **return** the sliding button to the start.
- **Remove** the detachable tube and **clean** the outside with distilled water.
- **Twist & remove** the cartridge from the Titrator and **attach** the cap to the cartridge. However, the cartridge could be left in the Titrator overnight in a secure location, if it is going to be used the next day.

Section 5: AT THE END OF THE FIELD WORK

1. Hanna Multiparameter (HI98494)

- immerse the **pH/ORP** electrode in '**electrode cleaning solution**' for 30 minutes.
- clean** the electrode with distilled water and put 3-4 drops of '**electrode storage solution**' in the cap and place the electrode (sensor) in the cap.
- when the use of the multiparameter is complete, **place** a little storage solution (1cm) in the storage container to keep all the sensors moist and screw the storage container on to the multiparameter.
- do not leave any water in the main storage container.
- remove** batteries, if it is going to be stored for long time (3+ months).
- clean** the exterior of the meter with a tissue wetted with water or 70% ethanol (preferred).

2. Geopacks Flowmeter

- clean** and dry the handheld unit, cables and impellor.
- remove** batteries from the control unit and store.

3. Valeport Flowmeter

- clean** with water or 70% ethanol (preferred), and dry, especially the handheld unit, cables and impellor.
- remove** batteries from the control unit and store.

4. Geotech pump

- check** that all parts are in the case.
 - clean** the equipment with a wet tissue for dust deposits and clean the case, if necessary.
- NB.** If the pump is being returned to UCAM, the battery should be left securely stored in Peru.

5. Titrator

- clean** with a wet tissue to remove any dust.
- check** that the cartridges are capped and that the caps are fixed tightly.
- place** the indicator pillows in the appropriate bags with the correct labels.

6. HDPE bottles

- clean** the HDPE bottles by soaking in 1% HCl acid for 2-4 hours, if available.
- and/or **wash** with plenty of distilled water until pH7.
- dry** and store away from sunlight and dust for the next field work.

7. 'Inventory sheet' in the 'Check list' Excel file

-complete the 'Inventory sheet' in the 'Check list' Excel file.

-include the quantities and quality of each piece of equipment and chemical.

-note any items that need replacing and the quantities needed in the right hand column, for planning the next field visit.

8. Bottles and tubes for labs

-prepare bottles and tubes for sending to the appropriate labs by separating them according to the relevant tests and site numbers.

Section 6: GLOSSARY OF TERMS

1. Water Quality Parameters

***Alkalinity:** the ability of water to absorb H^+ . In other words, it is a measure of water's capacity to neutralize acids = acid neutralizing capacity. Without this acid-neutralizing capacity, any acid added to a stream would cause an immediate change in the pH.

Alkalinity is the sum of the molar equivalent of the anions that will react with H^+ minus the molar equivalent of H^+ already present in the water. Generally, this equates to the amount of carbonate and bicarbonate ions in the water and so carbonate alkalinity typically represents alkalinity.

Measuring alkalinity is important in determining a stream's ability to neutralize acidic pollution from rainfall or wastewater. It is one of the best measures of the sensitivity of the stream to acid inputs. It is measured in mEq/l of $CaCO_3$. High enough alkalinity is really important for the success of freshwater species. Most river water is between 100-200 mg/l $CaCO_3$.

***Chemical oxygen demand (COD):** indicative measure of the amount of oxygen consumed by reactions in a measured solution. The most common application of COD is in quantifying the amount of oxidizable pollutants which will reduce available oxygen and, consequently, the quality of water for aquatic life.

The COD concentrations in surface waters ranges from 20-200 mg/l depending on the level of contaminants. Treated sewage water is discharged in the range 75-100 mg/l.

It is usually expressed as mass of oxygen consumed over volume of solution in milligrams per litre (mg/l).

***Conductivity:** a measure of water's capability to pass electrical flow determined by the concentration of ions present. These conductive ions come from dissolved salts and inorganic materials such as alkalis, chlorides, sulphides and carbonate compounds.

Dissolved salts and other inorganic chemicals conduct electrical current. As the salinity increases the conductivity increases. Organic compounds, such as sugars, oils, and alcohols, do not form ions that conduct electricity. In most cases, conductivity is positively correlated with Total Dissolved Solids (TDS).

Conductivity is measured in micro-siemens per centimetre ($\mu S/cm$) and reported as conductivity at 25°C. Distilled water should be in the range 0.5-3.0 $\mu S/cm$. Freshwater streams range between 50 to 1500 $\mu S/cm$ but, ideally, should be in the range 150 to 500 $\mu S/cm$ to support diverse aquatic life. Typically drinking water is in the range of 200 - 800 $\mu S/cm$, and sea water is about 5 S/m (or 50,000 $\mu S/cm$).

***Dissolved oxygen (DO):** a complex measure of how much oxygen (O_2) is dissolved in water, expressed as either a percentage of a theoretical maximum or an absolute amount in mg/l. Oxygen enters water by diffusion from the atmosphere, aeration as it falls over rocks and waterfalls, and as a product of photosynthesis and winds. DO levels are determined by temperature, atmospheric pressure, salinity, and pH.

DO levels determine the suitability of water bodies for life - healthy river water should generally have dissolved oxygen concentrations above 6.5-8 mg/L and between about 80-120 %. If DO levels drop below 5 mg/l, aquatic life is put under stress and below 2mg/l produces hypoxia in fish. At high concentrations, super-saturated water (above 120%) is also unsuitable for life.

***Hardness:** is a measure of the mineral (inorganic) content of the water, principally, the amount of dissolved calcium and magnesium; the greater the amount, the harder the water. Generally, hard water reduces the toxicity of trace metals to aquatic life; some metal ions form insoluble precipitates and are not available to organisms.

Water hardness is usually expressed in milligrams per litre (mg/l) of dissolved calcium and magnesium carbonate expressed as equivalent of calcium carbonate and measured in mEq/l of CaCO_3 . An acceptable level range for water hardness is 100-300ppm. Hard water is not a significant health risk. It can cause mineral build-up in plumbing, fixtures, and water heaters, and poor performance of soaps and detergents.

***Oxygen Reduction Potential (ORP):** is a measure of the cleanliness of the water and its ability to break down contaminants.

ORP is a measure of the tendency of a chemical species to acquire electrons from or lose electrons to an electrode and thereby be reduced or oxidised, respectively. The redox potential is measured in volts (V), or millivolts (mV). Each chemical species has its own intrinsic redox potential; for example, the more positive the ORP, the greater the species' affinity for electrons and tendency to be reduced.

An ORP level at 650-750mV will kill pathogens instantly. At ORP levels below 500 mV pathogens are killed in an hour, or so. Drinking water usually has an ORP at 650+mV. A safe ORP level of for freshwater fish is generally considered to be 300+mV and below 150mv aquatic life will struggle.

***% (atmospheric) Oxygen:** atmospheric oxygen concentration is ~21%, but only ~1% oxygen in water.

Where air and water meet, this disparity causes oxygen molecules in the air to be directly absorbed into the water. The greater the movement of the water the greater the rate of absorption. Water also shows higher absorbance of oxygen at lower temperatures and greater pressures.

Although water molecules contain an oxygen atom, aquatic organisms living in natural waters need dissolved oxygen (DO) (See above).

***pH (power of hydrogen):** pH is a measure of how acidic/basic water is. The numerical value of pH is determined by the molar concentration of hydrogen ions (H^+). The higher the H^+ concentration, the lower the pH, and the higher the OH^- concentration, the higher the pH. At a neutral pH of 7, the concentration of both H^+ ions and OH^- ions is 10^{-7} M. Thus the ions H^+ and OH^- are always paired – as the concentration of one increases, the other will decrease; regardless of pH, the sum of the ions must always equal 10^{-14} M. In general, water with more free hydrogen ions has a pH lower than 7 and is considered acidic, whereas water that has more free hydroxyl ions with a pH greater than 7 is basic (alkaline). The normal range for pH in surface water systems is 6.5 to 8.5, and the pH range for groundwater is 6 to 8.5.

***Total Dissolved solids (TDS):** TDS is the total amount of mobile charged ions, including minerals, salts or metals dissolved in a given volume of water, expressed in units of mg per litre of water (mg/l), or parts per million (ppm). TDS comprise inorganic salts (principally calcium, magnesium, potassium, sodium, sulphates, bicarbonates and chlorides) and trace amounts of organic matter dissolved in water. The maximum TDS levels recommended by the WHO is 300ppm. The higher the TDS, the higher the conductivity and the lower the pH. If the water contains beneficial minerals, a TDS level of 150-250 ppm is optimal for life.

The TDS changes the mineral content of water (rivers), high levels of TDS often harm aquatic species. Also, dissolved salts can dehydrate the skin of aquatic animals, which can be fatal.

***Total Nitrogen (TN):** is the sum of nitrate (NO_3), nitrite (NO_2), organic nitrogen and ammonia (all expressed as N). The nitrogen cycle is the means by which atmospheric nitrogen becomes available to living organisms. Nitrogen is essential for life, but excess nitrogen causes eutrophication in natural water and a danger to life. Water quality data suggests that appropriate reference levels for TN range from 0.12 to 2.2 mg/l.

***Total Phosphate (TP):** is a measure of all forms of phosphorus, dissolved and particulate. TP is a good way to measure phosphorus in lakes because it includes both ortho-phosphate and the phosphorus in plant and animal fragments suspended in lake water. TP levels are more stable and an annual mean can indicate the water quality and trophic state of a lake. Phosphorus is usually considered the “limiting nutrient” in aquatic ecosystems, in other words, changes in phosphorous concentration have greater impacts on productivity than changes in other nutrient concentrations.

The natural levels of total phosphorus in water are generally less than 0.03 mg/l while the natural levels of phosphate usually range from 0.005 to 0.05 mg/l.

***Turbidity:** turbidity is a measure of the degree to which water loses its transparency due to the presence of suspended particulates. The more total suspended solids (TSS) in the water, the murkier it is and the higher its turbidity. Turbidity is another good measure of water quality; the higher the turbidity, the lower the water quality for both human consumption and aquatic life.

Turbidity is usually measured in nephelometric turbidity units (NTU). This is an optical measurement of the amount of light scattered by suspended materials, when a light is shone through a water sample. The higher the intensity of scattered light, the higher the turbidity. Typically, turbidity levels are highly variable by season, local geology, water flow and weather events. During a low-flow period, most rivers and lakes are fairly clear with a turbidity reading below 10 NTU.

***Water temperature:** a measure that determines which species thrive in any body of water. There is an inverse relationship between dissolved oxygen and temperature; warm water holds less dissolved oxygen than cold water, and very warm water may not hold sufficient DO for aquatic life. Some compounds are also more toxic to aquatic life at higher temperatures. Fluctuations in water temperature have a significant impact on aquatic life.

2. Water chemistry

***Anions & Cations:** an ion is an atom or molecule that has a net electrical charge. Since the charge of the electron (considered negative by convention) is equal and opposite to that of the proton (considered positive by convention), the net charge of an ion is non-zero due to its total number of electrons being unequal to its total number of protons.

A **cation** is a positively charged ion, with fewer electrons than protons, while an **anion** is negatively charged, with more electrons than protons. Cations and anions attract each other because of their opposite electric charges, and readily form ionic compounds.

The most abundant **cations** present in water are calcium (Ca), magnesium (Mg), sodium (Na), and potassium (K); the most abundant **anions** are bicarbonate (HCO_3^-), chloride (Cl), and sulphate (SO_4). The dominant dissolved ion must be greater than 50% of the total.

Water is a compound of hydrogen and oxygen that results from the combustion of hydrogen. It is an excellent solvent and, consequently, natural water is not chemically pure; containing dissolved minerals, salts and organic compounds and also disperse and colloid disperse substances, like gases, in varying concentrations and compositions. Water contains different proportions of the following ions, depending on the source of the water:

Cations

- Calcium [Ca^{2+}]
- Magnesium [Mg^{2+}]
- Sodium [Na^+]
- Potassium [K^+]
- Ammonium [NH_4^+]
- Iron [$\text{Fe}^{2+/3+}$]
- Manganese [Mn^{2+}]

Ca^{2+}	HCO_3^-
Mg^{2+}	Cl^-
Na^+	NO_3^-
K^+	SO_4^{2-}
NH_4^+	
CO_2	
SiO_2	
Organic	
Solids	

Anions

- Hydrogen carbonate [HCO_3^-]
- Chloride [Cl^-]
- Nitrate [NO_3^-]
- Sulphate [SO_4^{2-}]

Gases / Solids

- Carbon dioxide [CO_2]
- Oxygen [O_2]
- Silicate [SiO_2]

***Coliformes:** coliform bacteria contain the enzyme B-galactosidase and are a commonly used indicator of the sanitary quality of water. Coliforms can be found in the aquatic environment, in soil and on vegetation. While coliforms themselves are not normally the cause of serious illness, they are used to indicate that other pathogenic organisms of faecal origin may be present including disease-causing bacteria, viruses, or protozoa and many multi-cellular parasites.

Escherichia coli (*E.coli*) is a coliform with an incubation period of 12-72 hours and an optimal growth temperature of 37°C. Unlike the general coliform group, *E. coli* are almost always of faecal origin and their presence is thus an effective confirmation of faecal contamination. Most strains of *E. coli* are harmless, but some can cause serious illnesses in humans.

***Metals:** many different metals can be found in water. The most dangerous are 'heavy' metals, including arsenic, cadmium, chromium, copper, lead, mercury, and selenium.

Arsenic: occurs in an organic form, which is harmless to humans, and an inorganic form, which can have major health impacts.

Long-term exposure to inorganic arsenic is linked to higher risks of lung, bladder, skin, liver, kidney, nasal passages, and prostate cancer. Other effects include thickening and discoloration of the skin, nausea, stomach pain, vomiting, diarrhoea, etc.

Cadmium: is a naturally occurring metal found in low concentrations but in fairly high levels in sewage sludge.

It is a non-essential element for aquatic life which is, potentially, harmed by it. It has been shown to cause toxic effects to the kidneys, bone defects, high blood pressure, and have reproductive system impacts.

Chromium: doesn't occur naturally in elemental form, but chromium compounds can be found in water in trace amounts.

Trivalent chromium is an essential trace element for humans - it removes glucose from blood, and it also plays a vital role in fat metabolism. However, hexavalent chromium is extremely toxic and is known for its genotoxic carcinogens. It can cause allergic and asthmatic reactions, diarrhoea, stomach and intestinal bleedings, cramps, and liver and kidney damage.

Copper: commonly found in aquatic systems, originating from both natural sources such as geological deposits, volcanic activity, and the erosion of rocks and soils; and anthropogenic sources such as, mining activities, agriculture, sludge from sewage treatment works, and pesticides.

Copper is an essential nutrient at low concentrations, however, at higher concentrations copper is toxic to aquatic life. It can lead to adverse effects on survival, growth, reproduction, brain function, blood chemistry, and metabolism.

Lead: a toxic heavy metal, not usually found naturally in water.

Exposure to lead can lead to premature birth, delayed physical and mental development in babies and cause learning disabilities. Accumulation of lead in adults can cause high blood pressure and kidney problems and may also cause anaemia, strokes, damage to the nervous system, and cancer.

Mercury: a naturally occurring, highly toxic metal and potent neurotoxin.

Scientists have found high levels of mercury accumulation in a wide range of wildlife species, causing dangerous reproductive and neurological problems. It impacts the function and development of the central nervous system in both people and wildlife.

Mercury bioaccumulates, which means that it increases in concentration with each step up in the food chain. When people consume large amounts of fish that contain mercury, they may experience neurological and gastrointestinal problems.

Selenium: a naturally occurring element present in sedimentary rocks, phosphate deposits and soils.

Selenium can enter surface water through weathering and erosion, or it can also be released during mining related activities, and irrigation for agriculture.

In small amounts selenium is an essential element for animals, but at higher concentrations is toxic. It bioaccumulates in the aquatic food chain and chronic exposure in fish and aquatic invertebrates can cause larval deformity or mortality. Selenium is also toxic to bird species that consume aquatic organisms containing excessive levels.

***Pesticides:** chemicals designed to control pests (and weeds) and can contain sulphur, chlorine, nitrogen, phosphorus, and bromine as well as heavy metals such as copper, arsenic, lead, and mercury. Pesticides include all of the following: herbicides, insecticides (including insect growth regulators, termiticides, etc), insect repellents, antimicrobials, and fungicide. The most common of these are herbicides which account for approximately 80% of all pesticide use, designed to protect plants/crops from weeds, fungi and insects.

Water pollution is caused by the improper use of pesticides that introduces chemicals into water, changing its properties and posing a threat to human and aquatic life.

3. Hydrology

Endorheic drainage: a closed-system **drainage basin** that retains water and allows no outflow to other external bodies of water, such as rivers or oceans, but converges instead into lakes or swamps, permanent or seasonal, that equilibrate through evaporation.

Exorheic drainage: an open-system drainage basin in which water drains to the sea. That is, outside the drainage basin. Such basins present many complex elements and characteristics throughout the water cycle of their entire system.

***Lentic:** refers to standing, non-flowing water such as water in pools, ponds and ditches.

***Lotic:** refers to water that is flowing such as water in streams and rivers.

4. Geology

Carbon flux: the amount of carbon exchanged between Earth's carbon pools - the oceans, atmosphere, land, and living things - and is typically measured in units of gigatonnes of carbon per year (GtC/yr).

Cation exchange: ion exchange in which one cation (such as sodium or hydrogen) is substituted for one or more other cations (such as calcium and magnesium in hard water).

Chemical species: a chemical substance or ensemble, composed of chemically identical molecular entities that can explore the same set of molecular energy levels on a characteristic or delineated time scale.

Denudation: the processes of erosion, leaching and reducing the mainland due to removal of material from higher to lower altitudes like river valleys, lakes and seas with a permanent filling of low lands.

Dissolution: the dissolving of rocks such as rock-salt (halite), gypsum and limestone (including chalk) producing features such as caves, sinkholes and large springs, creating a landscape known as *karst*.

Dolomitic: a sedimentary carbonate rock that contains a high percentage of the mineral dolomite, $\text{CaMg}(\text{CO}_3)_2$.

Oxidation: the loss of electrons during a reaction by a molecule, atom or ion.

Redox: is a type of chemical reaction in which the oxidation states of atoms are changed; characterized by the actual or formal transfer of electrons between chemical species, most often with one species undergoing **oxidation** while another species undergoes **reduction**.

Reduction: a gain of electrons or when the oxidation state of an atom, molecule, or ion decreases.

DATA RECORDING SHEET / RECORD de DATOS						
Date / Fecha:			Time / Hora:			
Site Name / Nombre del sitio:			GPS: (S / S) - (W/O) -			
Site code / Código del sitio:			Elevation / Altitud (metros):			
Air pressure (Presión de la atmosfera) (mbar):						
Air temperature (Temperatura del aire) (°C):						
Water data / Datos del agua	1	2	3	4	Av.	
pH						
Water temperature / M.						
Temperatura del agua (°C) MT.						
Conductivity / Conductividad (µ/cm)						
TDS (ppm)						
ORP (mV)						
Dissolved Oxygen / Oxígeno Disuelto (M - Multiparameter) (ppm)						
Dissolved Oxygen / Oxígeno Disuelto (MT - Mettler Toledo) (ppm)						
Dissolved Oxygen / Oxígeno Disuelto (M - Multiparameter) (%)						
Dissolved Oxygen / Oxígeno Disuelto (MT -Mettler Toledo) (%)						
River data / datos del río	1	2	3	4	5	Av.
Width / Ancho (metros)						
Depth / Profundidad (metros) (GF)						
Depth / Profundidad (metros) (VF)						
Velocity / Velocidad (m/s) (GF)						
Velocity / Velocidad (m/s) (VF)						
St.deviation/Deviación estándar (VF)						
CSA (m²):	Discharge / Caudal (m³/s):					
Water colour / Color del agua:						

River bed sediment types / Tipos de sedimentos del lecho del río:	<i>Sand/Arena</i>	<i>Pebbles/Guijarros</i> <i>Rock & Boulders/Rocas</i>	<i>Cobbles/Piedras</i>
Principal sediment colour / Color principal de los sedimentos:			
Sediment samples / Muestras de sedimentos:		No. / Número	
River bed / Lecho del río			
River bank / Orilla			
Water samples / Muestras de agua:		Size / Tamaño (ml)	
Metals (Filtered) / Metales (Filtrado) **Add Nitric Acid **			
Aniones + Cationes (Filtered / Física Química (Filtrado))			
COD / DQO (Unfiltered / Sin filtrar) ** Add Sulphuric Acid **			
Coliformes (Unfiltered / Sin filtrar)			
Pesticides (Unfiltered / Sin filtrar)			
Other / Otros:			
Empty bottle weight (gr) / Peso de la botella vacío:	Bottle weight with filtered water (gr) / Peso de la botella con agua filtrado:	Nett weight (gr) / Peso net:	
Filter papers used / Filtros usados:			
Field Laboratory Tests / Pruebas del laboratorio del campo			
Alkalinity / Alkalinidad (mg/l CaCO ₃)	Result / Resultado		
Tester kit in field / en el campo			
Test strips / Tiras reactivas			
Titration / Titulación		0.16N o 1.6N	
Natural observations / Observaciones naturales: <i>algae / algas, plants / plantas, geology/geología,</i> Human impact observations / Observaciones del impacto humano: <i>land use / uso del terreno, contamination / contaminación,</i>			
Data collectors / Recolectores de datos	Person / Persona	Universidad	
River characteristics / Características del río		UCAM / UNAH	
Water characteristics / Características del agua		UCAM / UNAH	
Water samples / Muestras de agua		UCAM / UNAH	
Sediment samples / Muestras de sedimentos		UCAM / UNAH	
Photos / Fotos		UCAM / UNAH	
Drone footage / Imágenes con drones		UCAM / UNAH	

Section 8: RISK ASSESSMENTS

Risk assessments ensure that everyone involved in the fieldwork and investigation are aware of the risks associated with the procedures and guidance is given on how to minimize the risks. Please read ALL the individual risk assessments associated with field and laboratory activities before you start work.

This section covers risk assessments regarding water sampling, preservation and field tests only. General fieldwork risk assessments can be found as a separate document, ask the field coordinators.

List of risk assessments

NB. These are not available in a digital format.

WCR01: Calibration of equipment for pH, Electrical Conductivity, Dissolved Oxygen, ORP measurements.

WCR02: Use of *sodium thiosulphate* for preserving water samples for pesticide tests.

WCR03: Testing for nitrates in water samples using Merck Cell Test Kits.

WCR04: Testing for Alkalinity in water samples using a Hach titrator.

WBR01: Water sampling, preservation and testing.