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Monitoring the presence and effects of marine litter in Mediterranean MPAs: the Plastic Busters MPAs approach

**PREPARED BY** 

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



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#### **Document Information**

This document (Deliverable 5.2.1) is a compilation of all the protocols that should be applied in order to elaborate a comprehensive diagnosis of the presence and effects of marine litter in Mediterranean MPAs.

#### **Approvals**

Date	Partner
20/4/2022	M.C. Fossi/UNISI (Project Scientific Coordinator, Task Leader)
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## **1.** Introduction

### 1.1 Marine litter a lurking threat in Mediterranean MPAs

The Mediterranean Sea is one of the areas most affected by marine litter worldwide. Marine litter any persistent, manufactured or processed solid material- is found lying on the shores, as well as floating anywhere from the surface to the bottom of the sea. Even in pristine environments of the Mediterranean, such as coastal and marine protected areas (MPAs), marine litter is building up, threatening habitats and species. Impacts vary from entanglement and ingestion, to bioaccumulation and bio-magnification of toxic substances released from litter items, facilitation of introduction of invasive species, damages to benthic habitats, etc. MPA managers stand at the forefront of this issue, and admittedly they lack the tools, knowledge, and often the resources to effectively tackle it. As a result, the achievement of the conservation goals set is hampered.



Figure 1-1. Marine litter a lurking threat in Mediterranean MPAs (Photo © Th. Vlachogianni).

### 1.2 The Plastic Busters MPAs project in a nutshell

The 4-year-long Interreg Med Plastic Busters MPAs project aimed at contributing to biodiversity protection and preservation of natural ecosystems in pelagic and coastal marine protected areas (MPAs), by defining and implementing a harmonized approach against marine litter. The project entailed actions that addressed the entire management cycle of marine litter, from monitoring and assessment to prevention and mitigation, as well as actions to strengthen networking between and among pelagic and coastal MPAs.

Plastic Busters MPAs consolidated Mediterranean efforts against marine litter by:

- Assessing the impacts of marine litter on biodiversity in MPAs and identifying marine litter 'hotspot' areas;
- Defining and testing tailor-made marine litter surveillance, prevention and mitigation measures in MPAs;
- Developing a common framework of marine litter actions for Interreg Mediterranean regions towards the conservation of biodiversity in Mediterranean MPAs.

The Plastic Busters MPAs project deployed the multidisciplinary strategy and common framework of action developed within the Plastic Busters initiative led by the University of Siena and the Sustainable Development Solutions Network Mediterranean (SDSN Med). This initiative frames the priority actions needed to tackle marine litter in the Mediterranean basin and was labelled under the Union for the Mediterranean (UfM) in 2016, gathering the political support of 43 Euro-Mediterranean countries.

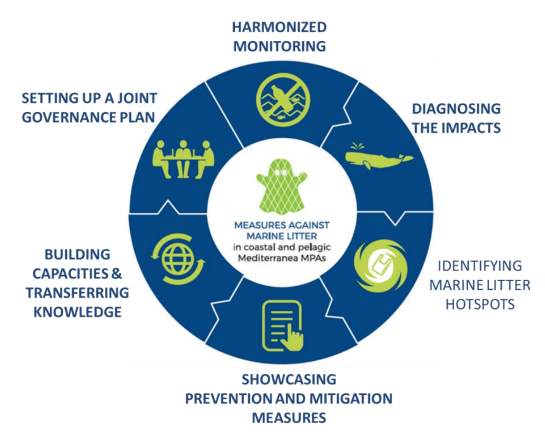


Figure 1-2. The Plastic Busters MPAs project in a nutshell.

### 1.3 Definitions and policy context

Within this document, marine litter is defined as any persistent, manufactured or processed solid material discarded, disposed of, or abandoned in the marine and coastal environment. Marine litter can be classified in size classes as follows: macrolitter refers to items larger than 25 mm in the longest dimension, mesolitter to items between 5 mm to 25 mm, and microlitter to items ranging from 1  $\mu$ m to 5 mm. This latter size class is sometime further broken down into large microlitter ranging from 1 mm to 5 mm and microplastic, from 1  $\mu$ m to 1 mm in size.

The main legislative frameworks related to marine litter monitoring are the EU Marine Strategy Framework Directive – MSFD (2008/56/EC, 2010/477/EC, 2017/848/EC) and the Barcelona Convention Ecosystem Approach (COP19 IMAP Decision IG.22/7, UNEP/MED WG.450/3, June 2018) (see Box 1.1 and Box 1.2).

**Box 1.1.** The Marine Litter Descriptor, criteria, and respective Indicators within the framework of the EU MSFD.

### Marine Litter within the EU MSFD

Descriptor 10: Properties and quantities of marine litter do not cause harm to the coastal and marine environment

**Criteria D10C1 - Primary:** The composition, amount and spatial distribution of litter on the coastline, in the surface layer of the water column, and on the seabed are at levels that do not cause harm to the coastal and marine environment.

- amount of litter washed ashore and/or deposited on coastlines, including analysis of its composition, spatial distribution and, where possible, source (10.1.1)
- amount of litter in the water column (including floating at the surface) and deposited on the seafloor, including analysis of its composition, spatial distribution and, where possible, source (10.1.2)

**Criteria D10C2 - Primary:** The composition, amount and spatial distribution of micro-litter on the coastline, in the surface layer of the water column, and in seabed sediment are at levels that do not cause harm to the coastal and marine environment.

 amount, distribution and, where possible, composition of microparticles (in particular microplastics) (10.1.3)

**Criteria D10C3 - Secondary:** The amount of litter and micro-litter ingested by marine animals is at a level that does not adversely affect the health of the species concerned.

• amount and composition of litter ingested by marine animals (10.2.1)

**Criteria D10C4 - Secondary:** The number of individuals of each species, which are adversely affected due to litter, such as by entanglement, other types of injury or mortality, or health effects.

**Box 1.2.** The Marine Litter Operational Objectives and respective Indicators within the framework of the Barcelona Convention Ecosystem Approach and the Integrated Monitoring and Assessment Programme (IMAP).

### Marine Litter and the Barcelona Convention Ecosystem Approach

**Ecological Objective 10 (EO10):** Marine and coastal litter do not adversely affect the coastal and marine environment.

### IMAP Common Indicator 22:

Trends in the amount of litter washed ashore and/or deposited on coastlines (including analysis of its composition, spatial distribution and, where possible, source).

### IMAP Common Indicator 23:

Trends in the amount of litter in the water column including micro plastics and on the seafloor.

#### IMAP Candidate Indicator 24:

Trends in the amount of litter ingested by, or entangling marine organisms, focusing on selected mammals, marine birds, and marine turtles.

### 1.4 About this document

The overarching aim of this document is to provide an operational protocol for implementing the Plastic Busters MPAs harmonized marine litter monitoring approach and assess the presence and effects of marine litter in pelagic and coastal Mediterranean MPAs with special emphasis on marine species, including endangered ones (cetaceans, sea turtles, birds, sharks, etc.). In this respect, this document is a compilation of all the protocols that should be applied in order to elaborate a comprehensive diagnosis of the marine litter problem in Mediterranean MPAs.

This document takes stock of all recent advances made by the EU MSFD Technical Group on Marine Litter and the Barcelona Convention CORMON Group. Furthermore, this document capitalizes on the outcomes of relevant projects such as the IPA-Adriatic DeFishGear project, the EU-funded INDICIT project and the Interreg Med marine litter related projects, namely the MEDSEALITTER, AMARE and ACT4LITTER.



Photo © Th. Vlachogianni



This document describes the methodological approach for monitoring macrolitter on beaches. It has been compiled based on the related methodology developed within the IPA-Adriatic DeFishGear project and the 2022 MSFD TGML Updated Guidance on Monitoring of Marine Litter in European Seas, while taking into account the results from the Plastic Busters MPAs testing phase.

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### 2.1. Site selection

The survey sites should fulfil the following characteristics:

- Have a minimum length of 100m;
- Be characterized by a low to moderate slope;
- Have clear access to the sea (not blocked by breakwaters or jetties);
- Be accessible to survey teams throughout the year;
- Ideally, not be subject to cleaning activities. In case they are subjected to litter collection activities, the timing of non-survey related beach cleaning must be known so that litter flux rates (the amount of litter accumulation per unit time) can be determined.

In addition, the location of the survey sites should be spatially stratified to reflect:

- different pressures and different levels of exposure to litter (e.g. close to river mouths, close to harbours/marinas, presence of touristic facilities nearby, etc.);
- different development and urbanisation levels, including a balanced mix of urban, semiurban, and remote/natural beaches.

It should be highlighted that all necessary precautions should be taken to ensure that surveys will not pose any threat to endangered or protected species such as sea turtles, shorebirds, marine mammals or sensitive beach vegetation/habitats.

### 2.2. Frequency and timing of surveys

At least four surveys should be carried out in winter, summer, spring and autumn. The optimum survey periods are:

- Winter: January
- Spring: April
- Summer: July
- Autumn: October

### 2.3. Sampling unit

A sampling unit is defined as a fixed section of a beach covering the whole area from the strandline to the back of the beach. The sampling unit should be a 100-metre stretch of beach along the strandline and reaching to the back of the beach. The back of the beach needs to be explicitly identified using coastal features such as the presence of vegetation, dunes, cliff base, road, fence or other anthropogenic structures such as seawalls (either piled boulders or concrete structures).

Sampling units should represent the general characteristics of the survey site and the general state of litter in the survey site. The sampling units should not be placed on the edges of a beach or on parts of the beach that have a higher potential to accumulate litter. In addition, the sampling unit should not be placed in potential litter hotspots such as areas near the entrance of the beach or near coastal parking lots or directly in front of hotels. Based on these considerations a set of potential sampling units should be identified and a random selection of sampling units should then be made from this set (e.g., dividing the coast into 100 m sections and randomly choosing a number of these sections as sampling units).

In case of heavily littered beaches, 100-metre stretches may be too difficult to survey and therefore two 50-metre stretches should be surveyed instead.

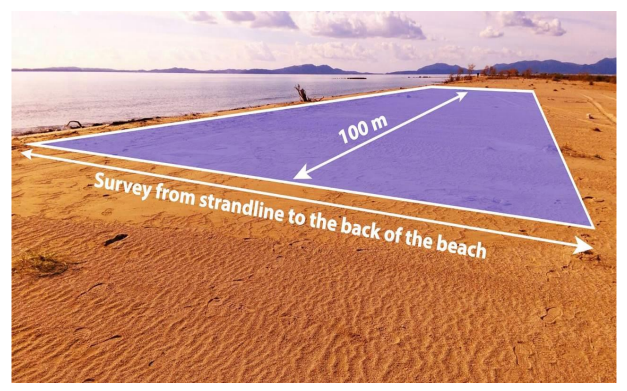


Figure 2-1. The sampling unit.



*Figure 2-2*. A heavily littered beach (Photo © Th. Vlachogianni).

### 2.4. Litter size classes to be surveyed

There are no upper size-limits for litter items to be recorded on beaches. But in order to ensure the inclusion of caps, lids, cigarette butts and other similar items in the quantification of beach litter, items as small as 2.5 cm in the longest dimension have to be recorded. In case such items are found in extremely high numbers, a 1-metre (rather than a 100-metre) beach transect should be used instead, saving effort and time.

### 2.5. Litter items classification and quantification

Items found on the sampling unit must be classified by type, according to the 'Joint List of Marine Litter Items Categories' prepared by the MSFD Technical Group on Marine Litter (MSFD TG ML) in close collaboration with EU Member States and the Regional Sea Conventions (Fleet et al., 2021). The manual for applying the Joint List classification system provides detailed information on how to classify litter items and a complementary photo guide helps the surveyors identify and categorise the litter items (<u>Online Photo Catalogue of the Joint List of Litter Categories</u>).

Litter items can be classified and recorded either on-site or in a working place (e.g. a lab) after the sampling has been completed (e.g., in case of bad weather conditions and/or heavily littered beaches); however, the latter should be avoided for weathered or fragile items, which easily disintegrate and can lead to overestimation of these litter items.

The unit to be used to assess the litter density is 'number of items' and should be expressed as counts of litter items per one 100-metre stretch.



Figure 2-3. Marine litter items classification (Photo © Th. Vlachogianni).

### 2.6. Litter items removal and disposal

During the survey, all litter items should be removed from the sampling unit. Larger items that cannot be removed (safely) by the surveyors should be marked, for example with paint spray (which meets environmentally friendly standards) so that they are not counted again at the next survey. The litter items collected should be disposed of properly. Regional or national regulations and arrangements should be followed. If these do not exist, local municipalities should be informed.

### 2.7. Materials and equipment

The following items are necessary to carry out beach surveys:

- High-resolution camera
- Hand-held GPS unit with extra batteries
- 100-metre tape measure (fiberglass preferred)
- Flag markers/stakes
- Rubbish bags
- Protective gloves
- Rigid container and sealable lid to collect sharp items such as needles, etc.
- Clipboard for each surveyor
- Recording sheets (printed on waterproof paper)
- Pencils and pens
- First aid kit (to include sunscreen, bug spray, drinking water)

### 2.8. Additional considerations

The amount and type of litter found on beaches can be influenced by different circumstances. To ensure that data will be analyzed and interpreted properly these circumstances must be recorded. Indicative examples of such circumstances include: events that may lead to unusual types and/or amounts of litter (e.g. shipping container losses, overflows of sewage treatment systems, etc.); difficult weather conditions (e.g. heavy winds or rain, etc.); replenishment/nourishment of the beach; etc.

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IPA-Adriatic DeFishGear, 2014. Methodology for Monitoring Marine Litter on Beaches (macro-debris >2.5 cm).

## Survey Site/Beach Identity Sheet

Nam	ne and area of survey site/beach:		
Bea	ch ID:		
Cou	Country:		
1.	Beach width at mean low spring tide (m):		
2.	Beach width at mean high spring tide (m):		
3.	Total length of the beach (m):		
4.	Back of beach (e.g. cliffs, dunes, etc.):		
5.	GPS coordinates of the four corners of the sampling unit:		
	A:		
	В:		
	C:		
	D:		
6.	Coordinate system used:		
7.	Date coordinates were measured://(d/m/y)		
8.	Prevailing sea currents off the beach: $\Box$ N $\Box$ E $\Box$ S $\Box$ W		
9.	Prevailing winds:  N E S W		
Whe	en you look from the beach to the sea, what direction is the beach facing: $\Box$ N $\Box$ E $\Box$ S $\Box$ W		
Type of beach material (% coverage):			

Prevalent beac	h usage (local people	, swimming and sunbathing, fishing, s	urfing, sailing, etc.):
1		seasonal or whole year round:	
2		seasonal or whole year round:	
3		seasonal or whole year round:	

Access to the beach:  $\Box$  Vehicle  $\Box$  Pedestrian  $\Box$  Boat\*

What is the distance to the What is the position of the survey area:	town in relation to the		
What is the (seasonal) popu			
Residential and tourist	winter	🗌 Tourist	winter
	spring		spring
	summer		summer
	autumn		autumn
Is there any development I	pehind the beach:	□No □Yes,	please describe
Are there food and/or drin			□No □Yes
What is the distance from t	he survey area to the food	and/or drink outlet (kr	
Present all year round:			□Yes □No please specify:
Position of food and/or drin	nk outlet in relation to the	survey area:	□n □e□s □w
What is the distance from	the beach to the nearest s	hipping lane (km):	
What is the estimated traff	ic density (number of ships	s/year) :	
Is it used mainly by mercha	nt ships, fishing vessels or	all kinds:	
Position of shipping lane in	relation to survey area:		□n □e□s □w
What is the distance from	the beach to the nearest h	arbor (km):	
What is the name of the ha	rbor:		
Position of harbor in relation	n to survey area:		□n □e□s □w
Type of harbor:			
Size of harbor (number of s	hips):		

What is the distance fro	m the beach to the nearest rive	r mouth (km):
What is the name of the	river:	
Position of river mouth i	n relation to survey area:	$\Box$ n $\Box$ e $\Box$ s $\Box$ w
Is the beach located nea	r a discharge or discharges of w	vaste water:
	n the beach to the discharge po	ints (km):
Position of discharge poi	nts in relation to survey area:	
How often is the beach	cleaned:	Daily Weekly Monthly Other
All year round:		
Seasonal, please specify	in months:	
		Daily Weekly Monthly Other
What method is used:		Manual Mechanical
Who is responsible for the	ne cleaning:	
Additional comments ar	nd observations about this beac	h:
Please include:		
1. A map of the beach		
2. A map of the beach ar following:	nd the local surroundings. When	relevant please mark on this map the
Nearest town	Food/drink outlets	Nearest shipping lane
Nearest harbor	Nearest river mouth	□ Discharge or discharges of waste water
3. A regional map		
Date sheet is filled in:	// (d/m/y)	
Name:		
E-mail:		

## Survey Sheet (100m)

e-mail address:		
Name of surveyor 2:		
e-mail address:		
Start time of the survey: End time of the survey:		
Information		
ect the data of the survey? If so, please tick appro		
🗌 Wind 🗌 Rain 🗌 Snow 🗌 Ice 🔲 Fog		
□ Sand storm □ Exceptionally high tide		
□ Yes □ No If so, how many: ame if known:		
🗆 Alive 🗆 Dead		

Were there any events that led to unusual types and/or amounts of litter on the beach? (For example

beach party or other)

Please specify: .....

## Joint List of Marine Macrolitter Items

## \* To be recorded also if smaller than 2.5 cm

J-CODE	SUP/FG	NAME	ITEMS COUNT
		ARTIFICIAL POLYMER MATERIALS	
J220		plastic sheeting from greenhouses	
J221		plastic irrigation pipes	
J222		other plastic items from agriculture	
J90		plastic flower pots	
J223		trays for seedlings of foamed plastic	
J46	FG	plastic oyster trays	
J45	FG	plastic mussels/oyster mesh bags, net sack, socks	
J47	FG	plastic sheeting from mussel culture (Tahitians)	
J102		plastic flip-flops	
J136		footwear made of plastic - not flip flops	
J40		plastic gloves (household/dishwashing, gardening)	
J41		plastic gloves (industrial/professional applications)	
J252		single-use plastic gloves	
J69		plastic hard hats/helmets	
J256		foamed plastic insulation including spray foam	
J89		plastic construction waste (not foamed insulation)	
J8	SUP	plastic drink bottles >0.5 l	
J7	SUP	plastic drink bottles ≤ 0.5 l	
J224	SUP	plastic food containers made of foamed polystyrene	
J21*	SUP	plastic caps/lids drinks	
J225	SUP	plastic food containers made of hard non-foamed plastic	
J1	SUP	plastic 4/6-pack yokes & six-pack rings	
J226	SUP	cups and cup lids of foamed polystyrene	
J227	SUP	cups and lids of hard plastic	
J228	SUP	plastic cutlery	
J229	SUP	plastic plates and trays	
J230	SUP	plastic stirrers	
J231	SUP	plastic straws	
J30	SUP	plastic crisps packets/sweets wrappers	
J31	SUP	plastic lolly & ice-cream sticks	
J85	FG	plastic commercial salt packaging	
J58	FG	fish boxes - foamed polystyrene	
J57	FG	fish boxes - hard plastic	
J92	FG	plastic bait containers/packaging	
J60*	FG	plastic fishing light sticks / fishing glow sticks incl. packaging	
J62	FG	plastic floats for fishing nets	
J59	FG	plastic fishing line	
J54	FG	plastic nets and pieces of net > 50cm	
J53	FG	plastic nets and pieces of net 2.5 cm $\geq \leq 50$ cm	
J232	FG	plastic string and filaments exclusively from dolly ropes	
J233	FG	other plastic string and filaments exclusively from fishery	
J234	FG	plastic tangled nets and rope without dolly rope or mixed with dolly rope	

J-CODE	SUP/FG	NAME	ITEMS COUNT
J235	FG	plastic tangled dolly rope	
J61	FG	other plastic fisheries related items not covered by	
101	гu	other categories	
J42	FG	plastic crab/lobster traps (pots) and tops	
J44	FG	plastic octopus pots	
J70		plastic shotgun cartridges	
J11		plastic beach use related body care and cosmetic bottles and containers	
J12		plastic non-beach use related body care and cosmetic bottles and containers	
J95	SUP	plastic cotton bud sticks	
J29		plastic combs/hair brushes/sunglasses	
J98		plastic diapers/nappies	
J236		other plastic personal hygiene and care items	
J96	SUP	plastic sanitary towels/panty liners/backing strips	
J144	SUP	plastic tampons and tampon applicators	
J97		plastic toilet fresheners	
J237	SUP	plastic wet wipes	
J253		plastic single-use face-mask	
J211		other plastic medical items (swabs, bandaging, adhesive plasters etc.)	
J100*		plastic medical/ pharmaceuticals containers/tubes/ packaging	
J99		plastic syringes/needles	
19		plastic bottles and containers of cleaning products	
J15		plastic engine oil bottles & containers >50cm	
J14		plastic engine oil bottles & containers 2.5 cm $\geq \leq 50$ cm	
J17		plastic injection gun containers/cartridges	
J16		plastic jerry cans	
J22*		plastic caps/lids chemicals, detergents (non-food)	
J23*		plastic caps/lids unidentified	
J24*		plastic rings from bottle caps/lids	
J13		other plastic bottles & containers (drums)	
J3	SUP	plastic shopping/carrier/grocery bags	
J101	501	plastic dog/pet faeces bag	
J5	SUP	the part that remains from tear-off plastic bags	
J36		other plastic heavy-duty sacks	
J238		plastic mesh bags for vegetable, fruit and other products	
J4	SUP	small plastic bags	
	301	plastic biomass holder from sewage treatment plants	
J91*		and aquaculture	
J18		plastic crates, boxes, baskets	
J65		plastic buckets	
193		plastic cable ties	
J84		plastic CDs & DVDs	
J67		plastic sheets, industrial packaging, sheeting	
J64		plastic fenders	
J68		fibre glass items	
J63		plastic floats/buoys other source than fishing or not	

J-CODE	SUP/FG	NAME	ITEMS COUNT
		known	
J239		other foamed plastic items and fragments not made of foamed polystyrene	
J257*		foamed plastic packaging	
J83		fragments of foamed polystyrene > 50cm	
J82		fragments of foamed polystyrene 2.5 cm ≥ ≤ 50 cm	
J80		fragments of non-foamed plastic > 50cm	
J79		fragments of non-foamed plastic 2.5cm ≥ ≤ 50cm	
J240		other identifiable foamed plastic items	
J241		other identifiable non-foamed plastic items	
J166		plastic paint brushes	
J28		plastic pens and pen lids	
J49		plastic rope (diameter more than 1cm)	
J242		plastic string and cord (diameter less than 1cm) not from dolly ropes or unidentified	
J66		plastic strapping bands	
J43		plastic tags (fishing, shipping, farming and industry)	
J87		plastic masking/duct/packing tape	
J88		telephone	
J72		plastic traffic cones	
J86		plastic fin trees (from fins for scuba diving)	
J243		plastic remains of fireworks	
J32*		plastic toys and party poppers	
J27*	SUP	tobacco products with filters (cigarette butts with filters)	
J26		plastic cigarette lighters	
J25		plastic tobacco pouches / plastic cigarette packet packaging	
J19		plastic vehicle parts	
		RUBBER	
J127		rubber boots	
J133		rubber condoms (incl. packaging)	
J131*		rubber band (small, for kitchen/household/post use)	
J248		rubber sheet	
J134		other rubber pieces	
J249		rubber belts	
J125*	SUP	rubber balloons	
J126		rubber balls	
J250		rubber inner-tubes	
J251		rubber tyres	
		CLOTH/TEXTILE	
J137		clothing	
J138		shoes & sandals made of leather and/or textile	
J141		cloth textile carpet & furnishing	
J140		hessian sacks/packaging	
J143		sails, canvas	
J145		other textiles	
J139		cloth textile backpacks & textile bags	

	PAPER/CARDBOARD	
J150	paper cartons/Tetrapak milk	
J151	paper cartons/Tetrapak (non-milk)	
J244	paper cups	
J245	paper food trays, food wrappers, drink containers	
J246	paper cotton bud sticks	
J240	other paper containers	
J147	paper bags	
J148	cardboard boxes	
J156	paper fragments	
J150 J154	paper newspapers & magazines	
J154 J158	other paper items	
J155	paper tubes and other pieces of fireworks	
J155 J152	paper tubes and other pieces of meworks paper cigarette packets	
J152	PROCESSED/WORKED WOOD	
J159	wooden corks	
	wooden ice-cream sticks, chip forks, chopsticks,	
J165	toothpicks	
J164	wooden fish boxes	
J163	wooden crab/lobster pots	
J162	wooden crates, boxes, baskets for packaging	
J172	other processed wooden items > 50cm	
J171	other processed wooden items 2.5 cm $\ge$ 50 cm	
J160	wooden pallets	
J167	wooden fireworks & matches	
	METAL	
J194	metal cables	
J175	metal drinks cans	
J176	metal food cans	
J181	metal tableware (e.g. plates, cups & cutlery)	
J184	metal lobster/crab pots	
J182*	metal fisheries related weights/sinkers, and lures	
J180	metal appliances (refrigerators, washers, etc.)	
J187	metal drums & barrels	
J174	metal aerosol/spray cans	
J188	other metal cans	
J190	metal paint tins	
J178*	metal bottle caps, lids & pull tabs from cans	
J195*	metal household batteries	
J177	metal foil wrappers, aluminium foil	
J199	other metal pieces > 50cm	
J198	other metal pieces 2.5cm $\geq \leq 50$ cm	
J186	metal industrial scrap	
J191	wire, wire mesh, barbed wire	
J179	metal disposable BBQs	
J193	metal vehicle parts / batteries	
J130	wheels with metal hub	

	GLASS/CERAMICS	
J204	glass ceramic construction materials (bricks, tiles, cement)	
J203	glass and ceramic tableware (plates/cups/glasses)	
J207	ceramic or glass octopus pots	
J200	glass bottles	
J201	glass jars	
J208	pieces of glass/ceramic (glass or ceramic fragments ≥ 2.5 cm)	
J205	glass fluorescent light tube	
J202	glass light bulbs	
J219	other ceramic items	
J210	other glass items	
	CHEMICALS	
J216	unidentified generally dark-coloured oil-like chemicals	
J217	unidentified generally light-coloured paraffin-like chemicals	
J218	unidentified chemicals	
	FOOD WASTE	
J215	organic food waste	



This document describes the methodological approach for monitoring microlitter on beaches. It has been compiled based on the related methodologies developed within the IPA-Adriatic DeFishGear project and the JPI-Oceans BASEMAN project, while taking into account the results from the Plastic Busters MPAs testing phase.

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### 3.1. Survey site selection

The survey sites for monitoring microlitter on beaches should be selected in accordance with the selection criteria of the survey sites for monitoring macrolitter; thus, the survey sites should fulfill the following characteristics:

- Have a minimum length of 100m;
- Be characterized by a low to moderate slope;
- Have clear access to the sea (not blocked by breakwaters or jetties);
- Be accessible to survey teams throughout the year;
- Ideally, not be subject to cleaning activities. In case they are subjected to litter collection activities, the timing of non-survey related beach cleaning must be known so that litter flux rates (the amount of litter accumulation per unit time) can be determined.

In addition, the location of the survey sites should be spatially stratified to reflect:

- different pressures and different levels of exposure to litter (e.g. close to river mouths, close to harbours/marinas, presence of touristic facilities nearby, etc.);
- different development and urbanisation levels, including a balanced mix of urban, semiurban, and remote/natural beaches.



Figure 3-1. Microlitter on beach sediment (Photo © Th. Vlachogianni).

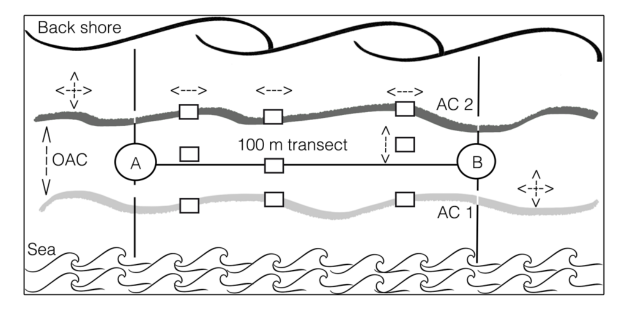
### 3.2. Frequency and timing of surveys

At least four surveys should be carried out in winter, summer, spring and autumn, at the same time with the beach macrolitter surveys. The optimum survey periods are:

- Winter: January
- Spring: April
- Summer: July
- Autumn: October

### 3.3. Sampling unit

The sampling area should be defined by marking out a 100-metre transect in width, parallel to the strandline, using a measuring tape and taking note of the GPS coordinates on each side of the transect (Fig. 3.2, A and B). The transect will define the sampling area i.e. from the shoreline (low tide, AC1) to above the strandline (accumulation zone, AC2). It should be highlighted that in many beaches the second tideline might not be always visible on the shore. Depending on the width of the beach, the sampling area can be extended to the back of the beach.



*Figure 3-2. Example of 100-metre transect (Frias et al., 2018) (AC: accumulation area, OAC: outside the accumulation area).* 

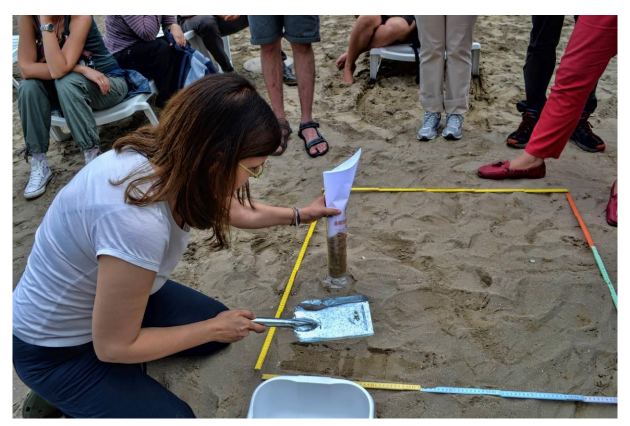


Figure 3-3. Microplastics sampling on beaches (Photo © Th. Vlachogianni).

A minimum of three samples along three transects vertical to the high tide line should be collected and the area between the two high tidelines should be surveyed. The sampling unit ( $30 \times 30 \text{ cm}$  or  $50 \times 50 \text{ cm}$  or  $1 \times 1 \text{ m}$ ) should be marked using a measuring tape or a quadrat and the GPS coordinates of each unit should be recorded. The top 3-5 cm of sediment should be sampled using a metal shovel or similar.

Large microplastics (1-5 mm) can be separated by sieving the beach sediment samples *in situ* through two metallic sieves with 1mm and 5mm mesh size; this is an effective method of reducing the sample volume. During sieving, the large or non-plastic items (e.g. shells, leaves, twigs, etc) should be removed. If the beach sediments are wet and difficult to go through the 1-mm sieve, the samples should be stored in glass jars or zip-lock bags and taken to the laboratory. The sediment samples should then be dried in the oven and then subsequently sieved.

### 3.4. Microlitter size classes to be surveyed

Typically litter items that are larger than 5mm and smaller than 2.5cm are sampled in microlitter surveys on beaches, however, the mesolitter items (items larger than 5mm and smaller than 2.5cm) that have been retained on the 5mm sieve can be surveyed too.

### 3.5. Litter analysis and classification

Concerning the separation of microplastics from the beach sediment, sieving is implemented for large microplastics (1-5mm), while floatation is used for small microplastics (<1mm) due to density differences between plastic and sediment particles. The principle of density floatation is commonly employed to separate less dense plastic polymers from denser sediment particles, and a range of high-density salt solutions have been used to extract microplastics from coastal and marine sediments. The floatation of the small microplastics is a rather demanding procedure, which should be carried out in the laboratory under specific conditions to avoid air-born contamination. All steps of the microplastics analysis must be conducted using 100% cotton lab coats and precautions are to be taken to avoid cross-contamination (e.g. airborne fibres).



Figure 3-4. Microplastics identification (Photo © Th. Vlachogianni).

The visual identification and classification of microlitter items can be carried out directly or through a microscope. Microplastics are characterized by **type** on the basis of the following categories: pellet, fragment (granule, flake), fibre, film, filaments, microbeads, foam (expanded polystyrene-PS). The most common **colours** of microplastics identified are the following: black, blue, white, transparent, red, green, multicolour, other. For the identification of the **polymer type** it is recommended to use an ATR-FTIR spectrometer or Raman spectroscopy.

### 3.6. Reporting units

Reporting units are extremely important to allow comparison among studies. The proposed reporting units for microplastics retrieved from sediment samples are:

- no. MPs per area (# particles m<sup>-2</sup>)
- no. MPs per volume (# particles m<sup>-3</sup>)
- no. MPs per mass (# particles kg<sup>-1</sup> dry sediment). In this case the weight of the sediment sample is needed or the density of the sediment
- mass of MP per area (g MP m<sup>-2</sup>)
- mass of MP per volume (g MP cm<sup>-3</sup>)

### 3.7. Materials and equipment

- High resolution camera
- Hand-held GPS unit, including extra batteries
- 100-metre tape measure (fiberglass preferred)
- Flag markers/stakes
- Metal shovel
- Metallic sieves (1mm and 5mm)
- Glass jars and paper bags
- Tweezers
- Recording sheets
- Pencils and pens
- First aid kit (to include sunscreen, bug spray, drinking water)
- Microscope
- ATR-FTIR spectrometer or Raman spectrometer

### References

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2)

Frias et al., 2018. Standardized protocol for monitoring microplastics in sediments. JPI-Oceans BASEMAN project.

GESAMP, 2019. Guidelines or the monitoring and assessment of plastic litter and microplastics in the ocean (Kershaw P.J., Turra A. and Galgani F. editors), (IMO/FAO/UNESCO-IOC/UNIDO/WMO/IAEA/UN/UNEP/UNDP/ISA Joint Group of Experts on the Scientific Aspects of Marine Environmental Protection). Rep. Stud. GESAMP No. 99, 130p.

IPA-Adriatic DeFishGear, 2014. Methodology for Monitoring Marine Microlitter on Beaches.

### 3.8. Survey sheets

An example of a datasheet is given below (from Frias et al, 2018). Alternatively, the survey datasheet used for the macrolitter surveys can be used, where the coordinates of each square should be also reported.

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d any of the following atmospheri	Fog   Smog	Dust- or san	d-storm	Waves exceptionally high
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C2 <u>1.</u> 2 AC 1 2				
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MPLING				
hore				
				1. Draw the high tide lines
			2	representing the main accumulation areas
				(AC1 and AC2);
				2. Mark starting point A and f
				point B. These should have 1 of distance between them;
				3. Draw the squares where
rnn	Land	22	2	sampling was conducted. (For example see Fig. 1).
<u> ~~~~~~</u>	SZZ	ひん	2	(, e. example see right)
ents/Notes:				



This document describes the methodological approach for monitoring identifying marine litter hotspots on beaches. It has been compiled based on the related methodology piloted within the Interreg Med AMARE project and it has been tested and adapted within the framework of the Plastic Busters MPAs project to address the recent advances in the field.

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### 4.1. Introduction

The present document presents a proposed approach to identify marine litter hotspots in Mediterranean MPAs using a simple protocol, which may also provide valuable insights for carrying out more comprehensive monitoring surveys using the protocols described in chapters 2 and 3. In addition, this methodological approach can provide the evidence needed for guiding targeted cleanup operations. It should be stressed that this approach does not provided a detailed assessment of amounts, types, composition and sources of marine litter, but it rather provides initial information on sites of interest.

The fate of most items is unknown and accumulations may occur at some locations as determined by several factors including hydrodynamic currents and circulation patterns, coastline structure, weather conditions, associated beach morphodynamics, residual swell, marine litter sources, both land-based and sea based. The amounts of litter observed thus reflect the long-term balance between inputs (land-based and sea-based sources and stranding processes) and removal (through export, burial, degradation and clean-ups). Apart from episodic storms events that may affect the number of items rather than the location of stranded items, most of the factors affecting the location of litter remain fairly constant with accumulation areas being the consequence of the integration of long term processes.

### 4.2. Methodological approach

Data are obtained from small boats (5-6 m) operating at low speed (1-12 knots) and moving at a distance of 20-100 m from the shore. The position of accumulation areas is recorded using GPS for low accumulation zones (2-10 litter items/site, usually a 5-30 m stretch distance onshore) and high accumulation zones (> than 10 litter items/site). The mapping of the hot spots of stranded litter is done through google maps, for simple analysis, or through a GIS (\*.shp) files mapping system, calculating the number of high accumulation areas ( > 10 items / site) load on 2 km or 3 km stretches of coastline. Maps are finally interpreted to support both the identification of potential monitoring sites, association with modelling predictions or identification of priority areas for removal actions.



**Figure 4-1.** (A) Visual observations and mapping of low (2-10 litter items/site, white circles) or high (> 10 litter items/site, red triangles) litter accumulation zones around the Elbe Island (Blue circles); (B) Mapping of high (> 10 litter items/site, red triangles) litter accumulation zones around the Elbe to locate priority areas for monitoring or cleaning.

## References

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

GESAMP (2019). Guidelines or the monitoring and assessment of plastic litter and microplastics in the ocean (Kershaw P.J., Turra A. and Galgani F. editors)). Rep. Stud. GESAMP No. 99, 130p.

5. Methodology for monitoring MACROLITTER on the sea-surface with visual observation by smalland medium-sized vessels

This document describes the methodological approach for monitoring macrolitter on the sea surface. It has been compiled based on the related methodologies developed within the IPA-Adriatic DeFishGear, the Interreg Med MEDSEALITTER projects and the 2022 MSFD TGML Updated Guidance on Monitoring of Marine Litter in European Seas, while taking into account the results from the Plastic Busters MPAs testing phase.

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### 5.1. Site selection

The monitoring of floating marine macrolitter by human observers is a methodology indicated for transects in selected areas. The selected areas should include:

- Low density areas (e.g. open sea);
- High density areas (e.g. close to ports);
- Other selected areas e.g. in estuaries, in the vicinity of cities, in local areas of touristic, recreational or commercial traffic.

Incoming currents from neighbouring areas or outgoing currents should be considered.



Figure 5-1. Floating macrolitter (Photo © Th. Vlachogianni).

### 5.2. Frequency and timing of surveys

At least two survey campaigns, one in autumn and one in spring should be carried out. The proposed campaign periods are:

- Autumn: October
- Spring: April

### 5.3. Sampling unit and sample size

The survey area is defined by the transect width and length. The transect width recommended to be used for small-scale vessels is 3 m on each side of the boat (6 m in total if two observers are deployed) and for medium-scale vessels 5 m on each side of the boat (10 m in total if two observers are deployed). The transect length should correspond approximately to 1 h of observation for each survey with a boat speed of 4-6 knots.

There is no agreed minimum sampling effort for obtaining a representative sample size and representative area coverage per survey campaign that can be extrapolated to all regions and/or density of litter situations for offshore and coastal waters, however for a transect width of 10 m, 15-30 h of effort have been recommended for monitoring an adequate sample size, while for a for a transect width of 6 m, 34-56 h of effort have been recommended (Aguilar et al., 2019).

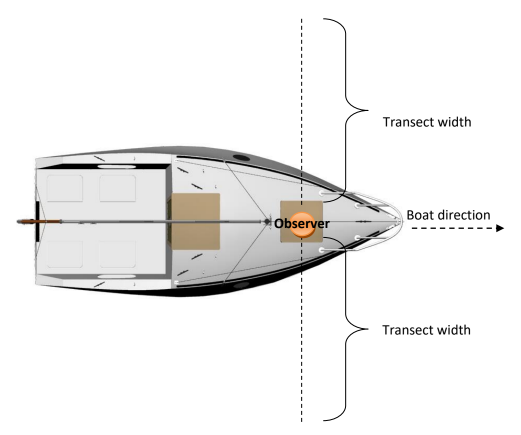


Figure 5-2. Schematic representation of observation position and transect width on a vessel.

### 5.4. Visual observation process

The observation should be made with naked eyes and binoculars can be used to confirm the litter sightings. A GPS is used to record the track of the monitored transect, to mark the beginning and the end of transect and indicate the position of the sighted objects. A telescopic fishing rod should be used in order to set the strip width.

The observation transect width should be set at 6 to 10 meters for small-sized vessels and mediumsized vessels respectively while the speed of the boat should not be higher than 4-6 knots. The observation, quantification and identification of floating litter items must be made by two dedicated observers who do not have other duties at the same time. The transect length should correspond approximately to 1 h of observation for each survey. The ideal location for observation is often the bow area of the boat. The observation direction must be perpendicular to the boat track (see figure below). The surveyors should conduct the survey from the glare-free side of the vessel and avoid the hours of the day when the sun is low on the horizon (sunrise and sunset), since visibility is not good enough due to glare and/or reflection. The surveys should be performed with sea state smaller or equal to 2 at the Beaufort scale.

#### 5.5. Litter size classes to be surveyed

Litter items larger than 2.5 cm (in the longest dimension) should be monitored and reported. Given that visual observation will not permit the exact measuring of object sizes, the following size range classes should be reported for each recorded litter item:

- A. 2.5 cm-5 cm
- B. 5 cm-10 cm
- C. 10 cm-20 cm
- D. 20 cm-30 cm
- E. 30 cm-50 cm
- F. 50 cm 100 cm
- G. >100 cm

#### 5.6. Litter classification and quantification

All items observed on the survey area should be classified by type, according to the 'Joint List of Marine Litter Items Categories' prepared by the MSFD Technical Group on Marine Litter (MSFD TG ML) in close collaboration with EU Member States and the Regional Sea Conventions (Fleet et al., 2021). The manual for applying the Joint List classification system provides detailed information on how to classify litter items and a complementary photo guide helps the surveyors identify and categorise the litter items (<u>Online Photo Catalogue of the Joint List of Litter Categories</u>). Data should be entered on the sheet while being observed.

Unknown litter or items that are not on the survey sheet should be noted in the appropriate "other item" category. A short description of the item should then be included on the survey sheet. If possible, digital photos should be taken of unknown items so that they can be identified later and, if necessary, be added to the survey sheet.

Furthermore, the occurrence of groups of floating litter items should be recorded along with their location as these could provide useful information with regards to accumulation areas. Ideally, each item in the group should be identified and recorded.

The unit in which litter will be assessed on the sea surface will be 'number of items' and it will be expressed as counts of litter items per square kilometer (litter items/km<sup>2</sup>). In order to compute the exact surveyed area, GPS coordinates must be recorded regularly (every min) to obtain an accurate measurement of the travelled transect. A handheld GPS unit might be handy in this respect.

#### 5.7. Materials and equipment

The following items are necessary to carry out floating litter surveys:

- Telescopic fishing rod;
- Digital camera;
- Binoculars;
- Hand-held GPS unit;
- Extra batteries (ideally rechargeable batteries);
- Clipboard for the surveyor;
- Recording sheets (printed on waterproof paper);
- Pencils;
- First aid kit (to include sunscreen, bug spray, drinking water).

## References

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# Monitoring MACROLITTER on the Water Surface Data Sheet

Location name	
Location ID	
Country	
Surveyor Name	
e-mail address	
Date of survey	

VESSEL CHARACTERISTICS				
Vessel name         Name of the vessel				
Type of vessel			Type e.g. research, fishing, hired, regular ferry etc.	
Vessel length and weight			Length of the vessel (metres) Gross weight of the vessel (tonnes)	

VISUAL SURVEY TRANSECT DETAILS				
Latitude/longitude start	Recorded as nnn.nnnn degrees at the start of the sample unit			
Latitude/longitude end	Recorded as nnn.nnnn degrees at the end of the sample unit			
Coordinates system	Datum and coordinate system employed			
Vessel speed	Average ship speed in knots			
Observation height	Observation elevation above the sea			
Distance covered	Total distance covered by the transect (m)			
Time start/end	Time over which the survey took place			
Surface covered	Area covered by the vessel (km <sup>2</sup> )			

ENVIRONMENTAL PARAMETERS - OBSERVATION DETAILS				
Wind speed		Recorded in (Beaufort)		
Wind direction		Tick more than one boxes e.g. for SE wind		
Sea surface salinity		Expressed in $^{0}/_{00}$ when reporting		
Viewing quality		Good/Moderate/Poor ; in the latter two case state cause (e.g. fog)		
Sea state		Expressed in accordance with the Douglas Sea Scale (0-9)		
NOTES				

SITE CHARACTERISTICS			-
Nearest river name			Name of nearest river
Nearest river distance			Distance to the nearest natural input (river or stream) (kilometers)
Nearest river position		□w	Position of river mouth in relation to survey area
Nearest major fishery			Name of the nearest major fishery (named by type)
Nearest major fishery distance			Distance to the nearest major fishery (kilometers)
Nearest major fishery position		□w	Position of the nearest major fishery in relation to survey area
Nearest town			Name of nearest town
Nearest town distance			Distance to the nearest town (kilometers)
Nearest town position		□w	Position of the nearest town in relation to survey area
Population size of this town			No of inhabitants
Additional features of the town	<ul> <li>Residential</li> <li>Tourist</li> <li>Residential &amp; tourist</li> </ul>	<ul> <li>Winter</li> <li>Spring</li> <li>Summer</li> <li>Autumn</li> </ul>	Indicate the main characteristic of the town, residential or touristic town; in case of the later indicate the high season peak
Name of the nearest beach			Name of the nearest beach
Distance to nearest beach			Distance to the closest coastline (kilometers)
Position of the nearest coast		□w	Position of the closest coastline in relation to survey area
Nearest shipping lane distance			Distance to the nearest shipping lane (kilometers)
Estimated traffic density			Recorded in number of ships/year
Vessel type			Indicate the type of vessels that mainly use it e.g. merchant ships, etc.
Position of the shipping lane		□w	Position of shipping lane in relation to survey area
Name of the nearest harbor			Name of nearest harbor
Distance to nearest harbor			Distance to the closest harbor (kilometers)
Harbor position		□w	Position of the nearest harbor in relation to survey area
Type of harbor			Based on the types of vessels visiting the harbor
Size of harbor			Record the number of ships that reach the harbor per year
Nearest discharge of waste water distance			Distance to the closest waste water discharge point(kilometers)
Position of nearest discharge point		□w	Position of nearest discharge points in relation to survey area
Type of waste water discharge	□Industrial □Munici	pal 🗌 Other	Indicate type of waste water discharged

## Joint List of Marine Macrolitter Items

\* To be recorded also if smaller than 2.5 cm

J-CODE	SUP/FG	NAME	ITEMS COUNT
		ARTIFICIAL POLYMER MATERIALS	
J220		plastic sheeting from greenhouses	
J221		plastic irrigation pipes	
J222		other plastic items from agriculture	
J90		plastic flower pots	
J223		trays for seedlings of foamed plastic	
J46	FG	plastic oyster trays	
J45	FG	plastic mussels/oyster mesh bags, net sack, socks	
J47	FG	plastic sheeting from mussel culture (Tahitians)	
J102		plastic flip-flops	
J136		footwear made of plastic - not flip flops	
J40		plastic gloves (household/dishwashing, gardening)	
J41		plastic gloves (industrial/professional applications)	
J252		single-use plastic gloves	
J69		plastic hard hats/helmets	
J256		foamed plastic insulation including spray foam	
J89		plastic construction waste (not foamed insulation)	
J8	SUP	plastic drink bottles >0.5 l	
J7	SUP	plastic drink bottles $\leq 0.5$ l	
J224	SUP	plastic food containers made of foamed polystyrene	
J21*	SUP	plastic caps/lids drinks	
		plastic food containers made of hard non-foamed	
J225	SUP	plastic	
J1	SUP	plastic 4/6-pack yokes & six-pack rings	
J226	SUP	cups and cup lids of foamed polystyrene	
J227	SUP	cups and lids of hard plastic	
J228	SUP	plastic cutlery	
J229	SUP	plastic plates and trays	
J230	SUP	plastic stirrers	
J231	SUP	plastic straws	
J30	SUP	plastic crisps packets/sweets wrappers	
J31	SUP	plastic lolly & ice-cream sticks	
J85	FG	plastic commercial salt packaging	
J58	FG	fish boxes - foamed polystyrene	
J57	FG	fish boxes - hard plastic	
J92	FG	plastic bait containers/packaging	
J60*	FG	plastic fishing light sticks / fishing glow sticks incl. packaging	
J62	FG	plastic floats for fishing nets	
J59	FG	plastic fishing line	
J54	FG	plastic nets and pieces of net > 50cm	
J53	FG	plastic nets and pieces of net 2.5 cm $\geq \leq 50$ cm	
J232	FG	plastic string and filaments exclusively from dolly ropes	
J233	FG	other plastic string and filaments exclusively from fishery	
J234	FG	plastic tangled nets and rope without dolly rope or mixed with dolly rope	

J-CODE	SUP/FG	NAME	ITEMS COUNT
J235	FG	plastic tangled dolly rope	
J61	FG	other plastic fisheries related items not covered by	
101	10	other categories	
J42	FG	plastic crab/lobster traps (pots) and tops	
J44	FG	plastic octopus pots	
J70		plastic shotgun cartridges	
J11		plastic beach use related body care and cosmetic bottles and containers	
J12		plastic non-beach use related body care and cosmetic bottles and containers	
J95	SUP	plastic cotton bud sticks	
J29		plastic combs/hair brushes/sunglasses	
J98		plastic diapers/nappies	
J236		other plastic personal hygiene and care items	
J96	SUP	plastic sanitary towels/panty liners/backing strips	
J144	SUP	plastic tampons and tampon applicators	
J97		plastic toilet fresheners	
J237	SUP	plastic wet wipes	
J253		plastic single-use face-mask	
J211		other plastic medical items (swabs, bandaging, adhesive plasters etc.)	
J100*		plastic medical/ pharmaceuticals containers/tubes/ packaging	
J99		plastic syringes/needles	
J9		plastic bottles and containers of cleaning products	
J15		plastic engine oil bottles & containers >50cm	
J14		plastic engine oil bottles & containers 2.5 cm $\geq \leq 50$ cm	
J17		plastic injection gun containers/cartridges	
J16		plastic jerry cans	
J22*		plastic caps/lids chemicals, detergents (non-food)	
J23*		plastic caps/lids unidentified	
J24*		plastic rings from bottle caps/lids	
J13		other plastic bottles & containers (drums)	
J3	SUP	plastic shopping/carrier/grocery bags	
J101		plastic dog/pet faeces bag	
J5	SUP	the part that remains from tear-off plastic bags	
J36		other plastic heavy-duty sacks	
J238		plastic mesh bags for vegetable, fruit and other products	
J4	SUP	small plastic bags	
J91*	501	plastic biomass holder from sewage treatment plants	
		and aquaculture	
J18		plastic crates, boxes, baskets	
J65		plastic buckets	
J93		plastic cable ties	
J84		plastic CDs & DVDs	
J67		plastic sheets, industrial packaging, sheeting	
J64		plastic fenders	
J68		fibre glass items	
J63		plastic floats/buoys other source than fishing or not	

J-CODE	SUP/FG	NAME	ITEMS COUNT
		known	
J239		other foamed plastic items and fragments not made of foamed polystyrene	
J257*		foamed plastic packaging	
J83		fragments of foamed polystyrene > 50cm	
J82		fragments of foamed polystyrene 2.5 cm ≥ ≤ 50 cm	
J80		fragments of non-foamed plastic > 50cm	
J79		fragments of non-foamed plastic 2.5cm ≥ ≤ 50cm	
J240		other identifiable foamed plastic items	
J241		other identifiable non-foamed plastic items	
J166		plastic paint brushes	
J28		plastic pens and pen lids	
J49		plastic rope (diameter more than 1cm)	
J242		plastic string and cord (diameter less than 1cm) not from dolly ropes or unidentified	
J66		plastic strapping bands	
J43		plastic tags (fishing, shipping, farming and industry)	
J87		plastic masking/duct/packing tape	
J88		telephone	
J72		plastic traffic cones	
J86		plastic fin trees (from fins for scuba diving)	
J243		plastic remains of fireworks	
J32*		plastic toys and party poppers	
J27*	SUP	tobacco products with filters (cigarette butts with filters)	
J26		plastic cigarette lighters	
J25		plastic tobacco pouches / plastic cigarette packet packaging	
J19		plastic vehicle parts	
		RUBBER	
J127		rubber boots	
J133		rubber condoms (incl. packaging)	
J131*		rubber band (small, for kitchen/household/post use)	
J248		rubber sheet	
J134		other rubber pieces	
J249		rubber belts	
J125*	SUP	rubber balloons	
J126		rubber balls	
J250		rubber inner-tubes	
J251		rubber tyres	
		CLOTH/TEXTILE	
J137		clothing	
J138		shoes & sandals made of leather and/or textile	
J141		cloth textile carpet & furnishing	
J140		hessian sacks/packaging	
J143		sails, canvas	
J145		other textiles	
J139		cloth textile backpacks & textile bags	

	PAPER/CARDBOARD	
J150	paper cartons/Tetrapak milk	
J151	paper cartons/Tetrapak (non-milk)	
J244	paper cups	
J245	paper food trays, food wrappers, drink containers	
J246	paper cotton bud sticks	
J247	other paper containers	
J147	paper bags	
J148	cardboard boxes	
J156	paper fragments	
J154	paper newspapers & magazines	
J158	other paper items	
J155	paper tubes and other pieces of fireworks	
J152	paper close and other precess of methods a paper close of the precess of methods and the precess of the preces	
5152	PROCESSED/WORKED WOOD	
J159	wooden corks	
J165	wooden ice-cream sticks, chip forks, chopsticks, toothpicks	
J164	wooden fish boxes	
J163	wooden crab/lobster pots	
J162	wooden crates, boxes, baskets for packaging	
J172	other processed wooden items > 50cm	
J171	other processed wooden items 2.5 cm $\geq 50$ cm	
J160	wooden pallets	
J167	wooden fireworks & matches	
	METAL	
J194	metal cables	
J175	metal drinks cans	
J176	metal food cans	
J181	metal tableware (e.g. plates, cups & cutlery)	
J184	metal lobster/crab pots	
J182*	metal fisheries related weights/sinkers, and lures	
J180	metal appliances (refrigerators, washers, etc.)	
J187	metal drums & barrels	
J174	metal aerosol/spray cans	
J188	other metal cans	
J190	metal paint tins	
J178*	metal bottle caps, lids & pull tabs from cans	
J195*	metal household batteries	
J177	metal foil wrappers, aluminium foil	
J199	other metal pieces > 50cm	
J198	other metal pieces 2.5cm ≥ ≤ 50cm	
J186	metal industrial scrap	
J191	wire, wire mesh, barbed wire	
J179	metal disposable BBQs	
J193	metal vehicle parts / batteries	
J130	wheels with metal hub	

	GLASS/CERAMICS		
J204	glass ceramic construction materials (bricks, tiles, cement)		
J203	glass and ceramic tableware (plates/cups/glasses)		
J207	ceramic or glass octopus pots		
J200	glass bottles		
J201	glass jars		
J208	pieces of glass/ceramic (glass or ceramic fragments ≥ 2.5 cm)		
J205	glass fluorescent light tube		
J202	glass light bulbs		
J219	other ceramic items		
J210	other glass items		
	CHEMICALS		
J216	unidentified generally dark-coloured oil-like chemicals		
J217	unidentified generally light-coloured paraffin-like chemicals		
J218	unidentified chemicals		
FOOD WASTE			
J215	organic food waste		



This document describes the methodological approach for monitoring microlitter on the sea surface. It has been compiled based on the related methodologies developed within the IPA-Adriatic DeFishGear project and the 2022 MSFD TGML Updated Guidance on Monitoring of Marine Litter in European Seas, while taking into account the results from the Plastic Busters MPAs testing phase.

**PREPARED BY** 

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



#### 6.1. Site selection

Given the high heterogeneity of litter distribution, the criteria for the survey sites selection could have crucial effect on results. The selection of the monitoring sites depends on the purpose and the methodology of monitoring, and can be made on the basis of certain characteristics of interest (i.e. MPAs of different scale such as large, medium or small) or through a random selection of survey sites.

For large scale MPAs, comprising of pelagic areas, sites of high and low microlitter accumulation should be surveyed.

<u>For medium and small scale MPAs</u>, confined to coastal waters around and in between small islands, an adequate number of sampling sites should be selected, based on the morphology and orientation of the island (shape, presence of inlets and gulfs, etc.) in order to cover all parts around the islands (N, S, E, W).

#### 6.2. Frequency and timing of surveys

At least two survey campaigns, one in autumn and one in early spring should be carried out. If possible, avoid periods with intense zooplankton blooms. The proposed campaign periods are:

- Autumn: October
- Spring: April

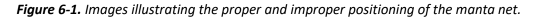
#### 6.3. Sampling unit and sample size

Manta nets are the most common tools for sampling floating mesolitter and microlitter in the surface layer of the water column. It is recommended that manta net should have mesh size of 333  $\mu$ m and be equipped with a flow meter. The sampling should be performed at low wind conditions (0-2 Beauforts) which can be recorded by a portable anemometer or by ship's instruments. The manta net should be towed for 30 min at a vessel speed that is maintained at less than 3 knots and both the start and end position should be recorded with GPS as well as the track.



(a) Correct

(b) Wrong





*Figure 6-2.* All tows should be conducted from the ship's side and beyond the ships' wake (Photo © Th. Vlachogianni).



*Figure 6-3.* After completion of each tow, the net should be washed thoroughly from the outside with filtered seawater (Photo © Th. Vlachogianni).

After completion of each tow, the net should be washed thoroughly from the outside with filtered seawater (<300 $\mu$ m) using the ship's hose in order to collect all particles in the cod-end. The sample collected in the cod-end should then be rinsed with seawater on a <300  $\mu$ m metallic sieve and transferred in glass jars with seawater. Any natural debris items, such as leaves, twigs, seaweed etc., should be rinsed separately above the sieve and removed from the sample. The samples should be stored in 70% ethanol solution for further analysis and a limited number of samples should be kept frozen to perform chemical analysis.



Figure 6-4. Sample on the sieve (a), cleared of any natural debris and tranferred in glass jars (b).



Figure 6-5. Sample rich in seaweeds before (a) and after (b) separation.

#### 6.4. Sample processing and size classification

The litter collected is classified in three size classes:

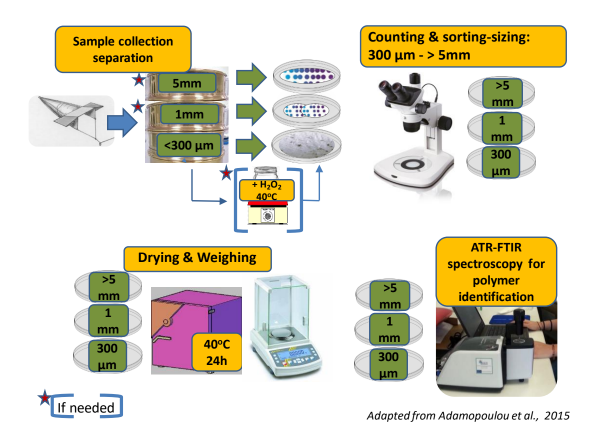
- Mesolitter (5 mm-25 mm)
- Large microlitter, LML (1mm-5mm)
- ▶ Small microlitter, SML (300µm 1mm)

In case of samples poor in natural particles and organic material, transfer the sample into a petri dish and observe under a stereomicroscope. Measure the particles' longest dimension using an image analysis software, count and classify into the 3 sizes classes. For the determination of weight, transfer the characterized MPs into three pre-weighted petri dishes according to size classes, dry at 40°C and weigh.

For samples rich in natural particles and organic material, the successive sieving as described below is helpful for the separation of the plastic particles but does not substitute the size characterization process with an image analysis software.

▶ Wet sieving and separation into 3 size classes: Pour the sample through a stacked arrangement of 5mm, 1mm, and 0.3 mm stainless steel mesh sieves. Accordingly, the litter items are classified in three size classes: small microlitter SML (300µm - 1mm), large microlitter LML (1mm-5mm), mesolitter (5 mm-25 mm).

- Mesolitter (5 mm-25 mm): Visually inspect the sample on the sieve, transfer and count only plastics in pre-weighted Petri dish. Dry at 40°C and weigh to determine the mass of mesoplastics.
- LML (1mm 5mm): Visually inspect the sample on the sieve, transfer and count the LML (1 mm-5 mm) particles in pre-weighed Petri dishes. Dry at 40°C and weigh to determine the mass of LMLs.
- SML (0.3mm -1mm): Collect the sample from the sieve with deionised water and filter through pre-weighed GF/C filters (pore size 1.2 μm). Dry the filters at 40°C for 24 hours and weigh. Determine the mass of small microlitter particles (SML mass). Examine the filter under a stereomicroscope and count SML particles.
- In case of high natural organic matter content in the samples (LML or SML) a step of peroxide digestion precedes filtration: Add 15% hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) with 1:1 (sample:H<sub>2</sub>O<sub>2</sub>) volume ratio and boil on a hot plate (approx.40°C) until the digestion is complete (no natural organic material should be visible). Collect the digested material with deionised water and continue with filtration, drying and mass determination.



*Figure 6-6.* Schematic representation of the various steps of processing floating mesolitter and microlitter samples.

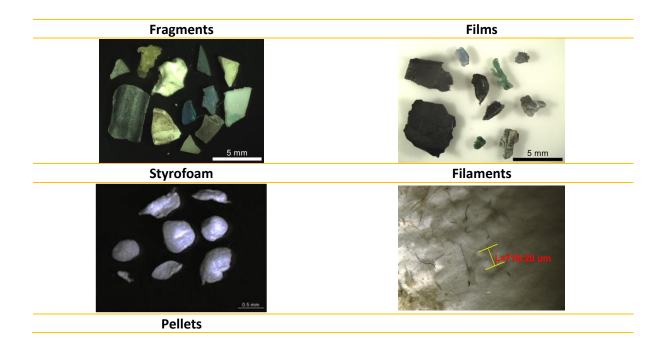
In all cases, for an accurate determination of the size of plastic particles, the particles collected in the petri dishes should be measured in their longest dimension under a stereoscope using an image analysis software, and then they should be classified in the three size classes (Meso, LMP, SMP). This is important because filaments and elongated particles may pass the sieves.



*Figure 6-7.* A microlitter sample (Photo © Th. Vlachogianni).

### 6.5. Sample analysis

Microplastics sorted, counted and characterized by **type** on the basis of the following categories: pellet, fragment (granule, flake), fibre, film, filaments, microbeads, foam (expanded polystyrene-PS), in line with the MSFD TGML guidelines. The most common **colours** of microplastics identified are the following: black, blue, white, transparent, red, green, multicolour, other. For the identification of the **polymer type** it is recommended to use an ATR-FTIR spectrometer or Raman spectroscopy or Pyrolysis-Gas chromatography-mass spectroscopy (Py-GCMS). FT-IR spectroscopy is mostly used in microplastic studies and in particular ATR-FTIR is considered fast, low cost and adequate for analyzing particles >300 µm, in size like the ones collected with manta nets.



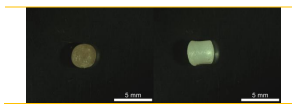


Figure 6-8. Examples of microlitter particles as seen under the stereomicroscope.

All surfaces should be clean. The glassware needs to be rinsed thoroughly with purified water. Samples should be covered with foil paper during the analysis. A glove bag should be used as working area for sample rinsing and filtrations. Petri dishes should be covered with glass lids during observation under stereomicroscope. Procedural blank samples should be used in all steps and items similar to those found in blank samples excluded, as they should be considered airborne contamination. Samples are to be kept in Petri dishes for long-term storage.

#### 6.6. Expression of the results

Microlitter counts (N) are reported as follows:

- N per km<sup>2</sup> or N per m<sup>2</sup>, based on the start end transect coordinates and the dimensions of the manta net mouth.
- N per Km<sup>3</sup> or N per m<sup>3</sup>, based on flow meter indication and relevant formula.

Microlitter mass is reported as follows:

- g per km<sup>2</sup> or g per m<sup>2</sup>
- g per Km<sup>3</sup> or g per m<sup>3</sup>

#### 6.7. Materials and equipment

#### Sampling equipment

- Manta net with wings and cod end (mesh size: 330 μm)
- Oceanographic flowmeter
- Submersible water pump with a hose (for rinsing the net) or other equipment for net rinsing
- GPS
- Glass jars with caps or plastic bottles (500 ml) (one or more per each sample; when on the sea is a lot of sea grass, than you need 2 - 3 plastic bottles per sample)
- Sample container cool box
- Screw driver
- Sieve (max 0.3 mm mesh size; preferable with smaller mesh size)
- ▶ Large bowl or washbasin (to prevent spillage of sample when emptying cod-end; 5 l <)
- Tap/fresh water source (tap/hose/squirt bottle)
- Squirt bottles2 x (one for water; one for alcohol)
- Tweezers (longer)
- Metal spoon
- Funnel (Ø 20 cm)
- Latex gloves without powder
- ▶ 70 % ethanol
- Waterproof marker, vellum paper and pencil

#### Sample separation equipment

- Stereomicroscope (min. 80x zoom; recommended also: transmission light with dark field, polarisation contrast and ring light)
- Object glasses (marked number of a sample, date of analysis)
- Micro tweezer and tweezer
- Glass petri dishes
- Lab coat
- ▶ 70 % ethanol
- 3 Sieves with mesh sizes: 5mm; 1mm; 0.3 mm or smaller
- Squirt bottle 2x (one for distilled water; one for alcohol)
- Latex gloves without powder
- Filtered water or distilled water
- Analytical laboratory scale
- Multiwell plate provided by NIC for storing the microlitter particles

# Monitoring MICROLITTER on the Water Surface Data Sheet

Location name	
Location ID	
Country	
Surveyor Name	
e-mail address	
Date of survey	

VESSEL CHARACTERISTICS					
Vessel name	Vessel name Name of the vessel				
Type of vessel			Type e.g. research, fishing, hired, regular ferry etc.		
Vessel length and weight			Length of the vessel (metres) Gross weight of the vessel (tonnes)		

MANTA NET TRANSECT DETAILS			
Latitude/longitude start	Recorded as nnn.nnnn degrees a the start of the sample unit		
Latitude/longitude end	Recorded as nnn.nnnn degrees a the end of the sample unit		
Coordinates system	Datum and coordinate system employed		
Vessel speed	Average ship speed in knots		
Distance covered	Total distance covered by the transect (m)		
Time start/end	Time over which the survey took place		

ENVIRONMENTAL PARAMETERS - OBSERVATION DETAILS			
Wind speed		Recorded in (Beaufort)	
Wind direction		Tick more than one boxes e.g. for SE wind	
Sea surface salinity		Expressed in % when reporting	
Viewing quality		Good/Moderate/Poor ; in the latter two case state cause (e.g. fog)	
Sea state		Expressed in accordance with the Douglas Sea Scale (0-9)	
NOTES.			

SITE CHARACTERISTICS			
Nearest river name			Name of nearest river
Nearest river distance			Distance to the nearest natural input (river or stream) (kilometers)
Nearest river position		Jw	Position of river mouth in relation to survey area
Nearest major fishery			Name of the nearest major fishery (named by type)
Nearest major fishery distance			Distance to the nearest major fishery (kilometers)
Nearest major fishery position		Jw	Position of the nearest major fishery in relation to survey area
Nearest town			Name of nearest town
Nearest town distance			Distance to the nearest town (kilometers)
Nearest town position		Jw	Position of the nearest town in relation to survey area
Population size of this town			No of inhabitants
Additional features of the town	<ul> <li>Residential</li> <li>Tourist</li> <li>Residential &amp; tourist</li> </ul>	<ul><li>Winter</li><li>Spring</li><li>Summer</li><li>Autumn</li></ul>	Indicate the main characteristic of the town, residential or touristic town; in case of the later indicate the high season peak
Name of the nearest beach			Name of the nearest beach
Distance to nearest beach			Distance to the closest coastline (kilometers)
Position of the nearest coast		Jw	Position of the closest coastline in relation to survey area
Nearest shipping lane distance			Distance to the nearest shipping lane (kilometers)
Estimated traffic density			Recorded in number of ships/year
Vessel type			Indicate the type of vessels that mainly use it e.g. merchant ships, etc.
Position of the shipping lane		Jw	Position of shipping lane in relation to survey area
Name of the nearest harbor			Name of nearest harbor
Distance to nearest harbor			Distance to the closest harbor (kilometers)
Harbor position		]w	Position of the nearest harbor in relation to survey area
Type of harbor			Based on the types of vessels visiting the harbor
Size of harbor			Record the number of ships that reach the harbor per year
Nearest discharge of waste water distance			Distance to the closest waste water discharge point(kilometers)
Position of nearest discharge point		Jw	Position of nearest discharge points in relation to survey area
Type of waste water discharge	□Industrial □Municipal □Other		Indicate type of waste water discharged



This document describes the methodological approach for monitoring macrolitter on the seafloor. It has been compiled based on the related methodology developed within the IPA-Adriatic DeFishGear and the 2022 MSFD TGML Updated Guidance on Monitoring of Marine Litter in European Seas, while taking into account the results from the Plastic Busters MPAs testing phase.

## PREPARED BY

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



#### 7.1. Site selection

Sites should be selected to ensure that they:

- ✓ Comprise areas with uniform substrate (ideally sand/silt bottom);
- ✓ Consider areas that might accumulate litter;
- ✓ Avoid areas of risk (presence of munitions), sensitive or protected areas;
- ✓ Do not exert impacts on any endangered or protected species.

Sites should be chosen following a two-fold approach: (i) selecting sites that meet certain criteria (e.g. are close to ports, river mouths, cities, etc.); (ii) choosing randomly from a large number of sites.

#### 7.2. Frequency and timing of surveys

The proposed survey periods are:

- Autumn: October
  - ✓ Spring: April

Following the MEDITS protocol, the hauls should be ideally performed during daylight. The daylight period is defined as the time between 30 minutes after sunrise and 30 minutes before sunset.

#### 7.3. Sampling unit and sample size

With regards to the sampling area, the MEDITS survey uses a depth stratified sampling scheme with random selection of trawling sites (same positions each year) within each stratum. Within this methodology, the following strata are proposed to be sampled: 10-50, 50-100, 100-200, 200-500 and 500-800 m. The size of the sampling area should be defined by each surveying team on the basis of the resources available for this task. The sampling density will be at least 2-3 stations (hauls) per 1000 km<sup>2</sup> in each stratum (average sampling density of the MEDITS survey) and/or 2-3 stations per sampling stratum.

#### 7.4. Trawling operation

Given that surveys might be performed with otter trawl fishing fleets or research vessels which use different gear (unlike the MEDITS surveys which use GOV nets), with no acoustic equipment, etc., it is evident that handling operations and parameters (such as type of mesh, mesh size of cod end, etc.) during the surveys cannot be standardized among and between the teams performing them. Nevertheless, MEDITS survey protocol should be followed as close as possible as described below:

- Haul position & orientation: The hauls should be positioned following a stratified sampling design, including at least three strata: 20-50 m; 51-100 m; 101-200 m, wherever possible. The hauls should be made over the same position in each sampling survey. The depth variations during the haul should not exceed ± 5% relative to the initial depth. The discrepancies to this target should be recorded. As far as possible, the hauls should be rectilinear.
- Haul speed & duration: The vessel speed should be 3 knots during the haul. However, if the skipper indicates that a slightly different speed is appropriate for optimal gear operation (depends on net characteristics) the vessel speed can be altered accordingly. In any case, vessel speed, hauling depth and geographical position should be continuously monitored during the haul (e.g. every 5 min). The haul duration is fixed at 30 min.
- Haul start and end definition: The start of the haul is defined as the moment at which the trawl geometry (vertical and horizontal) is stabilized. In the absence of electronic equipment

(acoustic devices like SCANMAR, etc.) the actual start time will be indicated by the skipper. The end of the haul is defined as the moment at which warp hauling begins.

• <u>Gear characteristics:</u> Cod-end mesh size and head rope length should be recorded.

#### 7.5. Litter size classes to be surveyed

The following size range classes will be reported for each recorded litter item:

- A.  $< 5 \text{ cm}^*5 \text{ cm} = 25 \text{ cm}^2$
- B. < 10 cm \* 10 cm = 100 cm  $^{2}$
- C. < 20 cm\*20 cm =  $400 \text{ cm}^2$
- D.  $< 50 \text{ cm}^* 50 \text{ cm} = 2500 \text{ cm}^2$
- E. < 100 cm-100 cm = 10000 cm<sup>2</sup> = 1 m<sup>2</sup>
- F. > 100 cm-100 cm = 10000 cm<sup>2</sup> = 1 m<sup>2</sup>

#### 7.6. Litter classification and quantification

All items collected from the haul must be classified by type, according to the 'Joint List of Marine Litter Items Categories' prepared by the MSFD Technical Group on Marine Litter (MSFD TG ML) in close collaboration with EU Member States and the Regional Sea Conventions (Fleet et al., 2021). The manual for applying the Joint List classification system provides detailed information on how to classify litter items and a complementary photo guide helps the surveyors identify and categorise the litter items (Online Photo Catalogue of the Joint List of Litter Categories).

Unknown litter or items that are not on the survey sheet should be noted in the appropriate "other item" box. A short description of the item should then be included on the survey sheet. If possible, digital photos should be taken of unknown items so that they can be identified later and, if necessary, be added to the survey sheet.

The unit in which litter should be recorded is the number of items and it should be expressed as counts of litter items per square kilometer (litter items/km<sup>2</sup>). The total weight of litter items per haul should be recorded, as well as the weight per each main litter category. In addition, the total weight of each haul should be recorded, as well as the weight of the commercial fish caught in it.

The estimation of litter items/km<sup>2</sup> requires the estimation of the "swept area". The latter is difficult to be monitored accurately during the haul because it requires the use of specialized equipment, like acoustic devices mounted on the trawl net. Such instruments might not be available during the samplings. However, the skipper, knowing by experience the geometry of the gear, can advise the surveying team on the effective mouth width and height of the net during each fishing operation.

Alternatively, the swept area can also be estimated following the method of Sparre and Venema (1998). The trawl sweeps a path, the area of which is the length of the path times the width of the trawl, called the "swept area" or the "effective path swept". The swept area (**a**) can be estimated by:

Where:

*V* is the velocity of the trawl over the ground when trawling;

*h* is the length of the head-rope;*D* is the cover of distance;

*t* is the time spent trawling;

**X** is that fraction of the head-rope length, which is equal to the width of the path swept by the trawl. The value of X varies from 0.4 to 0.66 for tropical waters and a value of X = 0.5 has been suggested as the best compromise value for the Mediterranean Sea (Sparre and Venema, 1992).

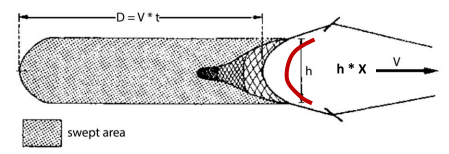


Figure 7-1. Swept area (source: <u>http://www.fao.org/docrep/w5449e/w5449e0f.htm</u>)

When exact positions of the start and the end of the haul are available the distance covered can be estimated in units of nautical miles (nm), by:

$$D = 60^* \sqrt{(Lat1 - Lat2)^2 + (Lor1 - Lor2)^2 * \cos^2(0.5^* (Lat1 + Lat2))}$$
.....

Where:

Lat1 = latitude at start of haul (degrees) Lat2 = latitude at end of haul (degrees) Lon1 = longitude at start of haul (degrees) Lon2 = longitude at end of haul (degrees)

#### 7.7. Materials and equipment

The following items are necessary to carry out beach surveys:

- Bucket or box
- Tape measure
- Plastic bags to collect the litter
- Digital camera
- Hand-held GPS unit, including extra batteries
- Digital scales for weighing litter (ideally with a 1g precision)
- Clipboard for the surveyor and recording sheets
- Pencils or pens
- First aid kit (to include sunscreen, bug spray, drinking water).

### Reference

Fleet, D., Vlachogianni, Th., Hanke, G., 2021. A Joint List of Litter Categories for Marine Macrolitter Monitoring. EUR 30348 EN, Publications Office of the European Union, Luxembourg, 2020, ISBN 978-92-76-21445-8, JRC121708. <u>https://doi.org/10.2760/127473</u>

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

IPA-Adriatic DeFishGear, 2014. Methodology for monitoring marine litter on the Seafloor with bottom trawl surveys.

# 7.8. Recording sheets

# Monitoring MACROLITTER on the Seafloor Data Sheet

Location name	
Location ID	
Country	
Surveyor Name	
e-mail address	
Date of survey	

VESSEL CHARACTERISTICS				
Vessel name	Name of the vessel			
Type of vessel	Type e.g. research, fishing, hired, regular ferry etc.			
Vessel length and tonnage	Length of the vessel (metres) Gross weight of the vessel (tonnes)			
Vessel engine power	Vessel engine power (kilowatt)			

HAUL DETAILS	
Latitude/longitude start	Recorded as nnn.nnnnn degrees at the start of the sample unit
Latitude/longitude end	Recorded as nnn.nnnnn degrees at the end of the sample unit
Coordinates system	Datum and coordinate system employed
Vessel speed	Average vessel speed in knots
Start time/end time	Time over which the survey (haul) took place
Mouth horizontal/vertical opening	Record the trawl mouth horizontal and vertical opening (mm)

Haul position/depth	Record the average haul posit	ion
Cod end mesh size	Record mesh size (mm)	
Cod end type	Type of cod end e.g. diamond mesh, square mesh	
Head rope length	Record the length of the head rope (m)	

ENVIRONMENTAL PARAMETERS - OBSERVATION DETAILS			
Wind speed			Recorded in (Beaufort)
Wind			Tick more than one boxes e.g. for SE wind
Sea state			Expressed in accordance with the Douglas Sea Scale (0-9)
<u>NOTES</u>			
SITE CHARACTERISTICS			
Nearest river name			Name of nearest river
Nearest river distance			Distance to the nearest natural input (river or stream) (kilometers)
Nearest river position	Nearest river position $\Box N \Box E \Box S \Box W$		Position of river mouth in relation to survey area
Nearest major fishery			Name of the nearest major fishery (named by type)
Nearest major fishery distance		Distance to the nearest major fishery (kilometers)	
Nearest major fishery position			Position of the nearest major fishery in relation to survey area
Nearest town			Name of nearest town
Nearest town distance			Distance to the nearest town (kilometers)
Nearest town position			Position of the nearest town in relation to survey area
Population size of this town			No of inhabitants
Additional features of the town	of the town Residential Winter Tourist Residential & tourist Autumn		Indicate the main characteristic of the town, residential or touristic town; in case of the later indicate the high season peak

Name of the nearest beach		Name of the nearest beach
Distance to nearest beach		Distance to the closest coastline (kilometers)
Position of the nearest coast		Position of the closest coastline in relation to survey area
Nearest shipping lane distance		Distance to the nearest shipping lane (kilometers)
Estimated traffic density		Recorded in number of ships/year
Vessel type		Indicate the type of vessels that mainly use it e.g. merchant ships, etc.
Position of the shipping lane	Ωn □e□s □w	Position of shipping lane in relation to survey area
Name of the nearest harbor		Name of nearest harbor
Harbor position	□n □e□s □w	Position of the nearest harbor in relation to survey area
Type of harbor		Based on the types of vessels visiting the harbor
Size of harbor		Record the number of ships that reach the harbor per year
Nearest discharge of waste		
water distance		
Position of nearest discharge		Position of nearest discharge points in
point		relation to survey area
NOTES		

### Joint List of Marine Macrolitter Items

## \* To be recorded also if smaller than 2.5 cm

J-CODE	SUP/FG	NAME	ITEMS COUNT
		ARTIFICIAL POLYMER MATERIALS	
J220		plastic sheeting from greenhouses	
J221		plastic irrigation pipes	
J222		other plastic items from agriculture	
J90		plastic flower pots	
J223		trays for seedlings of foamed plastic	
J46	FG	plastic oyster trays	
J45	FG	plastic mussels/oyster mesh bags, net sack, socks	
J47	FG	plastic sheeting from mussel culture (Tahitians)	
J102		plastic flip-flops	
J136		footwear made of plastic - not flip flops	
J40		plastic gloves (household/dishwashing, gardening)	
J41		plastic gloves (industrial/professional applications)	
J252		single-use plastic gloves	
J69		plastic hard hats/helmets	
J256		foamed plastic insulation including spray foam	
J89		plastic construction waste (not foamed insulation)	
J8	SUP	plastic drink bottles >0.5 l	
J7	SUP	plastic drink bottles ≤ 0.5 l	
J224	SUP	plastic food containers made of foamed polystyrene	
J21*	SUP	plastic caps/lids drinks	
J225	SUP	plastic food containers made of hard non-foamed plastic	
J1	SUP	plastic 4/6-pack yokes & six-pack rings	
J226	SUP	cups and cup lids of foamed polystyrene	
J227	SUP	cups and lids of hard plastic	
J228	SUP	plastic cutlery	
J229	SUP	plastic plates and trays	
J230	SUP	plastic stirrers	
J231	SUP	plastic straws	
J30	SUP	plastic crisps packets/sweets wrappers	
J31	SUP	plastic lolly & ice-cream sticks	
J85	FG	plastic commercial salt packaging	
J58	FG	fish boxes - foamed polystyrene	
J57	FG	fish boxes - hard plastic	
J92	FG	plastic bait containers/packaging	
J60*	FG	plastic fishing light sticks / fishing glow sticks incl. packaging	
J62	FG	plastic floats for fishing nets	
J59	FG	plastic fishing line	
J54	FG	plastic nets and pieces of net > 50cm	
J53	FG	plastic nets and pieces of net 2.5 cm $\geq \leq 50$ cm	
J232	FG	plastic string and filaments exclusively from dolly ropes	
J233	FG	other plastic string and filaments exclusively from	

J-CODE	SUP/FG	NAME	ITEMS COUNT
		fishery	
1224	50	plastic tangled nets and rope without dolly rope or	
J234	FG	mixed with dolly rope	
J235	FG	plastic tangled dolly rope	
J61	FG	other plastic fisheries related items not covered by	
	-	other categories	
J42	FG	plastic crab/lobster traps (pots) and tops	
J44	FG	plastic octopus pots	
J70		plastic shotgun cartridges	
J11		plastic beach use related body care and cosmetic bottles and containers	
J12		plastic non-beach use related body care and cosmetic bottles and containers	
J95	SUP	plastic cotton bud sticks	
J29		plastic combs/hair brushes/sunglasses	
J98		plastic diapers/nappies	
J236		other plastic personal hygiene and care items	
J96	SUP	plastic sanitary towels/panty liners/backing strips	
J144	SUP	plastic tampons and tampon applicators	
J97		plastic toilet fresheners	
J237	SUP	plastic wet wipes	
J253		plastic single-use face-mask	
J211		other plastic medical items (swabs, bandaging, adhesive plasters etc.)	
J100*		plastic medical/ pharmaceuticals containers/tubes/ packaging	
J99		plastic syringes/needles	
J9		plastic bottles and containers of cleaning products	
J15		plastic engine oil bottles & containers >50cm	
J14		plastic engine oil bottles & containers 2.5 cm $\ge \le 50$ cm	
J17		plastic injection gun containers/cartridges	
J16		plastic jerry cans	
J22*		plastic caps/lids chemicals, detergents (non-food)	
J23*		plastic caps/lids unidentified	
J24*		plastic rings from bottle caps/lids	
J13		other plastic bottles & containers (drums)	
J3	SUP	plastic shopping/carrier/grocery bags	
J101		plastic dog/pet faeces bag	
J5	SUP	the part that remains from tear-off plastic bags	
J36		other plastic heavy-duty sacks	
J238		plastic mesh bags for vegetable, fruit and other products	
J4	SUP	small plastic bags	
J91*		plastic biomass holder from sewage treatment plants and aquaculture	
J18		plastic crates, boxes, baskets	
J65		plastic buckets	
		•	

J-CODE	SUP/FG	NAME	ITEMS COUNT
J84		plastic CDs & DVDs	
J67		plastic sheets, industrial packaging, sheeting	
J64		plastic fenders	
J68		fibre glass items	
J63		plastic floats/buoys other source than fishing or not	
102		known	
J239		other foamed plastic items and fragments not made of	
J257*		foamed plottin polyaging	
		foamed plastic packaging	
J83		fragments of foamed polystyrene > 50cm	
J82		fragments of foamed polystyrene 2.5 cm $\geq \leq 50$ cm	
J80		fragments of non-foamed plastic > 50cm	
J79		fragments of non-foamed plastic 2.5cm $\geq \leq 50$ cm	
J240		other identifiable foamed plastic items	
J241		other identifiable non-foamed plastic items	
J166		plastic paint brushes	
J28		plastic pens and pen lids	
J49		plastic rope (diameter more than 1cm)	
J242		plastic string and cord (diameter less than 1cm) not from dolly ropes or unidentified	
J66		plastic strapping bands	
J43		plastic tags (fishing, shipping, farming and industry)	
J87		plastic masking/duct/packing tape	
J88		telephone	
J72		plastic traffic cones	
J86		plastic fin trees (from fins for scuba diving)	
J243		plastic remains of fireworks	
J32*		plastic toys and party poppers	
J27*	SUP	tobacco products with filters (cigarette butts with filters)	
J26		plastic cigarette lighters	
J25		plastic tobacco pouches / plastic cigarette packet	
		packaging	
J19		plastic vehicle parts	
		RUBBER	
J127		rubber boots	
J133		rubber condoms (incl. packaging)	
J131*		rubber band (small, for kitchen/household/post use)	
J248		rubber sheet	
J134		other rubber pieces	
J249		rubber belts	
J125*	SUP	rubber balloons	
J126		rubber balls	
J250		rubber inner-tubes	
J251		rubber tyres	
		CLOTH/TEXTILE	
J137		clothing	
J138		shoes & sandals made of leather and/or textile	

J-CODE	SUP/FG	NAME	ITEMS COUNT			
J141		cloth textile carpet & furnishing				
J140		hessian sacks/packaging				
J143		sails, canvas				
J145		other textiles				
J139		cloth textile backpacks & textile bags				
PAPER/CARDBOARD						
J150		paper cartons/Tetrapak milk				
J151		paper cartons/Tetrapak (non-milk)				
J244		paper cups				
J245		paper food trays, food wrappers, drink containers				
J246		paper cotton bud sticks				
J247		other paper containers				
J147		paper bags				
J148		cardboard boxes				
J156		paper fragments				
J154		paper newspapers & magazines				
J158		other paper items				
J155		paper tubes and other pieces of fireworks				
J152		paper cigarette packets				
		PROCESSED/WORKED WOOD				
J159		wooden corks				
J165		wooden ice-cream sticks, chip forks, chopsticks, toothpicks				
J164		wooden fish boxes				
J163		wooden crab/lobster pots				
J162		wooden crates, boxes, baskets for packaging				
J172		other processed wooden items > 50cm				
J171		other processed wooden items 2.5 cm $\ge$ 50 cm				
J160		wooden pallets				
J167		wooden fireworks & matches				
	1	METAL	1			
J194		metal cables				
J175		metal drinks cans				
J176		metal food cans				
J181		metal tableware (e.g. plates, cups & cutlery)				
J184		metal lobster/crab pots				
J182*		metal fisheries related weights/sinkers, and lures				
J180		metal appliances (refrigerators, washers, etc.)				
J187		metal drums & barrels				
J174		metal aerosol/spray cans				
J188		other metal cans				
J190		metal paint tins				
J178*		metal bottle caps, lids & pull tabs from cans				
J195*		metal household batteries				
J177		metal foil wrappers, aluminium foil				
J199		other metal pieces > 50cm				
J198		other metal pieces 2.5cm $\geq \leq$ 50cm				

J-CODE	SUP/FG	NAME	ITEMS COUNT			
J186		metal industrial scrap				
J191		wire, wire mesh, barbed wire				
J179		metal disposable BBQs				
J193		metal vehicle parts / batteries				
J130		wheels with metal hub				
GLASS/CERAMICS						
J204		glass ceramic construction materials (bricks, tiles, cement)				
J203		glass and ceramic tableware (plates/cups/glasses)				
J207		ceramic or glass octopus pots				
J200		glass bottles				
J201		glass jars				
J208		pieces of glass/ceramic (glass or ceramic fragments ≥ 2.5 cm)				
J205		glass fluorescent light tube				
J202		glass light bulbs				
J219		other ceramic items				
J210		other glass items				
CHEMICALS						
J216		unidentified generally dark-coloured oil-like chemicals				
J217		unidentified generally light-coloured paraffin-like chemicals				
J218		unidentified chemicals				
FOOD WASTE						
J215		organic food waste				

HAUL RESULTS	
Total weight of litter in the haul	Record litter weight in Kg
Total weight of artificial polymer materials	Record litter weight in Kg
Total No of items of artificial polymer materials	Record number of items
Total weight of rubber	Record litter weight in Kg
Total No of items of rubber	Record number of items
Total weight of cloth/textile	Record litter weight in Kg
Total No of items of cloth/textile	Record number of items
Total weight of paper/cardboard	Record litter weight in Kg
Total No of items of paper/cardboard	Record number of items
Total weight of processed/worked wood	Record litter weight in Kg
Total No of items of processed/worked wood	Record number of items
Total weight of metal	Record litter weight in Kg
Total No of items of metal	Record number of items
Total weight of glass/ceramics	Record litter weight in Kg
Total No of items of glass/ceramics	Record number of items



This document describes the methodological approach for monitoring macrolitter on the seafloor. It has been compiled based on the related methodology developed within the IPA-Adriatic DeFishGear and the 2022 MSFD TGML Updated Guidance on Monitoring of Marine Litter in European Seas, while taking into account the results from the Plastic Busters MPAs testing phase.

**PREPARED BY** 

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



#### 8.1. Site selection

Sites should be selected to ensure that they:

- Consider areas that might accumulate litter;
- Avoid areas of risk (presence of munitions and other hazardous waste), sensitive areas;
- Do not exert impacts on any endangered or protected species;
- Avoid areas with strong currents or waves;
- Avoid navigation routes of vessels that might put divers in danger.

Sites should be chosen following a two-fold approach: (i) selecting sites that meet certain criteria (e.g. are close to ports, river mouths, cities, etc.); (ii) choosing randomly from a large number of sites.

#### 8.2. Frequency and timing of surveys

At least two survey campaigns, one in autumn and one in spring should be carried out. The proposed survey periods are:

- Autumn: October
- Spring: April

If surveys are also implemented in the summer time these should be carried out in July.

#### 8.3. Sampling unit

The sampling unit is defined by the transect width and length. The line transects are defined with a nylon line, marked every 5 meters with resistant paints, that is deployed using a diving reel while SCUBA diving. Distances should be determined either by laying out a 100m tape measure or alternatively by laying a 100m length of weighted rope across the bottom. The start and end point of each transect should be identified with marker buoys and recorded using a GPS.

The length of the line transects could vary between 50m-100m and the width from 4m-8m, depending on the depth, the depth gradient, the turbidity, the habitat complexity and the litter density (see table below).

**Table 8-1**. Suggested transect lengths and widths of the sampling unit based on environmental conditions and litter densities (Katsavenakis, 2009).

Litter Density	Environmental Conditions	Sampling Unit (length x width)
0.1 – 1 items / m <sup>2</sup>	Low turbidity & high habitat complexity	20 m x 4 m
0.1 – 1 items / m <sup>2</sup>	High turbidity	20 m x 4 m
0.01 – 0.1 items / m <sup>2</sup>	In every case	100 m x 8 m
< 0.01 items / m <sup>2</sup>	In every case	200 m x 8 m

#### 8.4. Litter size classes to be surveyed

The following size range classes should be reported for each recorded litter item:

- A.  $< 5 \text{ cm}^*5 \text{ cm} = 25 \text{ cm}^2$
- B. < 10 cm \*10 cm = 100 cm  $^{2}$
- C.  $< 20 \text{ cm}^2 20 \text{ cm} = 400 \text{ cm}^2$
- D. < 50 cm\*50 cm = 2500 cm<sup>2</sup>
- E. < 100 cm\*100 cm = 10000 cm<sup>2</sup> = 1 m<sup>2</sup>
- F. > 100 cm\*100 cm = 10000 cm<sup>2</sup> = 1 m<sup>2</sup>

#### 8.5. Litter classification and quantification

All items collected from the sampling unit must be classified by type, according to the 'Joint List of Marine Litter Items Categories' prepared by the MSFD Technical Group on Marine Litter (MSFD TG ML) in close collaboration with EU Member States and the Regional Sea Conventions (Fleet et al., 2021). The manual for applying the Joint List classification system provides detailed information on how to classify litter items and a complementary photo guide helps the surveyors identify and categorise the litter items (Online Photo Catalogue of the Joint List of Litter Categories).

When conducting underwater visual surveys with a self-contained underwater breathing apparatus (scuba), lighter litter items should be collected (while larger items should just be marked), brought ashore and entered in the related seafloor litter monitoring sheet. When conducting underwater visual surveys with snorkelling, digital photos should be taken for all items with an underwater camera and subsequently should be entered in the seafloor litter monitoring sheet once identified. Unknown litter, or items that are not on the survey sheet, should be noted in the appropriate "other item" box. A short description of the items should then be included on the survey sheet.

The unit in which litter should be recorded is number of items and it should be expressed as counts of litter items per square kilometre (litter items/km<sup>2</sup>).



Figure 8-1. Litter items collected during a seafloor survey (Photo © Th. Vlachogianni).

#### 8.6. Materials and equipment

The following items are necessary to carry out seafloor litter surveys:

- Scuba gear and equipment: diving suit, buoyancy control device, regulator, air tank, compass, pressure gauge, fins, gloves, knife, and boots, etc.;
- Supplies: mesh sack, rope, ruler, cutter, dive flag, dive slate, float tube, and pelican float;
- Underwater digital camera;
- Lift bag;
- Floating fence;
- ► GPS;
- Comprehensive first-aid kit;
- Recording sheets and pencils.

### References

Fleet, D., Vlachogianni, Th., Hanke, G., 2021. A Joint List of Litter Categories for Marine Macrolitter Monitoring. EUR 30348 EN, Publications Office of the European Union, Luxembourg, 2020, ISBN 978-92-76-21445-8, JRC121708. <u>https://doi.org/10.2760/127473</u>

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IPA-Adriatic DeFishGear, 2014. Methodology for monitoring marine litter on the Seafloor-Visual surveys with SCUBA/snorkeling (Shallow coastal waters, 0–20m).

### 8.7. Sampling & recording sheets

### Monitoring Marine Litter (Macro) on the Seafloor Data Sheet

Location name	
Location ID	
Country	
Surveyor Name	
e-mail address	
Date of survey	

SITE DETAILS	
Latitude/longitude start	Recorded as nnn.nnnn degrees at the start of the sampling unit
Latitude/longitude end	Recorded as nnn.nnnn degrees at the end of the samplin unit
Length of sampling unit	Record length in m
Width of sampling unit	Record width in m
Depth	Record depth in m
Coordinates system	Datum and coordinate system employed
Start time/end time	Time over which the survey took place

ENVIRONMENTAL PARAMETERS - OBSERVATION DETAILS									
Underwater visibility	□Low □Moderate □High	Tick one box based on expert judgment							
Current velocity/turbidity	□Low □Moderate □High	Tick one box based on expert judgment							
Type of substrate	Rocky Sandy Mixed	Tick one box based on expert judgment							
Substrate complexity	□Low □Medium □High	Tick one box based on expert judgment							
Wind speed		Recorded in (Beaufort)							
Wind		Tick more than one boxes e.g. for SE wind							
Sea state		Expressed in accordance with the Douglas Sea Scale (0-9)							

SITE CHARACTERISTICS			
Nearest river name			Name of nearest river
Nearest river distance			Distance to the nearest natural input (river or stream) (kilometers)
Nearest river position		□w	Position of river mouth in relation to survey area
Nearest major fishery			Name of the nearest major fishery (named by type)
Nearest major fishery distance			Distance to the nearest major fishery (kilometers)
Nearest major fishery position		□w	Position of the nearest major fishery in relation to survey area
Nearest town			Name of nearest town
Nearest town distance			Distance to the nearest town (kilometers)
Nearest town position		□w	Position of the nearest town in relation to survey area
Population size of this town			No of inhabitants
Additional features of the town			Indicate the main characteristic of the town, residential or touristic town; in case of the later indicate the high season peak
Name of the nearest beach			Name of the nearest beach
Distance to nearest beach			Distance to the closest coastline (kilometers)
Position of the nearest coast		□w	Position of the closest coastline in relation to survey area
Nearest shipping lane distance			Distance to the nearest shipping lane (kilometers)
Estimated traffic density			Recorded in number of ships/year
Vessel type			Indicate the type of vessels e.g. merchant ships, etc.
Position of the shipping lane		□w	Position of shipping lane in relation to survey area
Name of the nearest harbor			Name of nearest harbor
Harbor position		□w	Position of the nearest harbor in relation to survey area
Type of harbor			Based on the types of vessels visiting the harbor
Size of harbor			Record the number of ships that reach the harbor per year
Nearest discharge of waste water distance			Name nearest location if waste water discharge
Position of nearest discharge point		□w	Position of nearest discharge points in relation to survey area

#### Joint List of Marine Macrolitter Items

### \* To be recorded also if smaller than 2.5 cm

J-CODE	SUP/FG	NAME	ITEMS COUNT
		ARTIFICIAL POLYMER MATERIALS	
J220		plastic sheeting from greenhouses	
J221		plastic irrigation pipes	
J222		other plastic items from agriculture	
J90		plastic flower pots	
J223		trays for seedlings of foamed plastic	
J46	FG	plastic oyster trays	
J45	FG	plastic mussels/oyster mesh bags, net sack, socks	
J47	FG	plastic sheeting from mussel culture (Tahitians)	
J102		plastic flip-flops	
J136		footwear made of plastic - not flip flops	
J40		plastic gloves (household/dishwashing, gardening)	
J41		plastic gloves (industrial/professional applications)	
J252		single-use plastic gloves	
J69		plastic hard hats/helmets	
J256		foamed plastic insulation including spray foam	
J89		plastic construction waste (not foamed insulation)	
J8	SUP	plastic drink bottles >0.5 l	
J7	SUP	plastic drink bottles ≤ 0.5 l	
J224	SUP	plastic food containers made of foamed polystyrene	
J21*	SUP	plastic caps/lids drinks	
J225	SUP	plastic food containers made of hard non-foamed plastic	
J1	SUP	plastic 4/6-pack yokes & six-pack rings	
J226	SUP	cups and cup lids of foamed polystyrene	
J227	SUP	cups and lids of hard plastic	
J228	SUP	plastic cutlery	
J229	SUP	plastic plates and trays	
J230	SUP	plastic stirrers	
J231	SUP	plastic straws	
J30	SUP	plastic crisps packets/sweets wrappers	
J31	SUP	plastic lolly & ice-cream sticks	
J85	FG	plastic commercial salt packaging	
J58	FG	fish boxes - foamed polystyrene	
J57	FG	fish boxes - hard plastic	
J92	FG	plastic bait containers/packaging	
J60*	FG	plastic fishing light sticks / fishing glow sticks incl. packaging	
J62	FG	plastic floats for fishing nets	
J59	FG	plastic fishing line	
J54	FG	plastic nets and pieces of net > 50cm	
J53	FG	plastic nets and pieces of net 2.5 cm $\geq \leq 50$ cm	
J232	FG	plastic string and filaments exclusively from dolly ropes	
J233	FG	other plastic string and filaments exclusively from	

J-CODE	SUP/FG	NAME	ITEMS COUNT
		fishery	
1224	ГC	plastic tangled nets and rope without dolly rope or	
J234	FG	mixed with dolly rope	
J235	FG	plastic tangled dolly rope	
J61	FG	other plastic fisheries related items not covered by	
	-	other categories	
J42	FG	plastic crab/lobster traps (pots) and tops	
J44	FG	plastic octopus pots	
J70		plastic shotgun cartridges	
J11		plastic beach use related body care and cosmetic bottles and containers	
J12		plastic non-beach use related body care and cosmetic bottles and containers	
J95	SUP	plastic cotton bud sticks	
J29		plastic combs/hair brushes/sunglasses	
J98		plastic diapers/nappies	
J236		other plastic personal hygiene and care items	
J96	SUP	plastic sanitary towels/panty liners/backing strips	
J144	SUP	plastic tampons and tampon applicators	
J97		plastic toilet fresheners	
J237	SUP	plastic wet wipes	
J253		plastic single-use face-mask	
J211		other plastic medical items (swabs, bandaging, adhesive plasters etc.)	
J100*		plastic medical/ pharmaceuticals containers/tubes/ packaging	
199		plastic syringes/needles	
19		plastic bottles and containers of cleaning products	
J15		plastic engine oil bottles & containers >50cm	
J14		plastic engine oil bottles & containers 2.5 cm $\ge \le 50$ cm	
J17		plastic injection gun containers/cartridges	
J16		plastic jerry cans	
J22*		plastic caps/lids chemicals, detergents (non-food)	
J23*		plastic caps/lids unidentified	
J24*		plastic rings from bottle caps/lids	
J13		other plastic bottles & containers (drums)	
J3	SUP	plastic shopping/carrier/grocery bags	
J101		plastic dog/pet faeces bag	
J5	SUP	the part that remains from tear-off plastic bags	
J36		other plastic heavy-duty sacks	
J238		plastic mesh bags for vegetable, fruit and other products	
J4	SUP	small plastic bags	
J91*		plastic biomass holder from sewage treatment plants and aquaculture	
J18		plastic crates, boxes, baskets	
J65		plastic buckets	
193		plastic cable ties	
122		plastic capie ties	

J-CODE	SUP/FG	NAME	ITEMS COUNT
J84		plastic CDs & DVDs	
J67		plastic sheets, industrial packaging, sheeting	
J64		plastic fenders	
J68		fibre glass items	
J63		plastic floats/buoys other source than fishing or not	
103		known	
J239		other foamed plastic items and fragments not made of	
1257*		foamed polystyrene	
J257*		foamed plastic packaging	
J83		fragments of foamed polystyrene > 50cm	
J82		fragments of foamed polystyrene 2.5 cm $\geq \leq 50$ cm	
J80		fragments of non-foamed plastic > 50cm	
J79		fragments of non-foamed plastic 2.5cm $\geq \leq 50$ cm	
J240		other identifiable foamed plastic items	
J241		other identifiable non-foamed plastic items	
J166		plastic paint brushes	
J28		plastic pens and pen lids	
J49		plastic rope (diameter more than 1cm)	
J242		plastic string and cord (diameter less than 1cm) not from dolly ropes or unidentified	
J66		plastic strapping bands	
J43		plastic tags (fishing, shipping, farming and industry)	
J87		plastic masking/duct/packing tape	
J88		telephone	
J72		plastic traffic cones	
J86		plastic fin trees (from fins for scuba diving)	
J243		plastic remains of fireworks	
J32*		plastic toys and party poppers	
J27*	SUP	tobacco products with filters (cigarette butts with filters)	
J26		plastic cigarette lighters	
J25		plastic tobacco pouches / plastic cigarette packet	
		packaging	
J19		plastic vehicle parts	
		RUBBER	
J127		rubber boots	
J133		rubber condoms (incl. packaging)	
J131*		rubber band (small, for kitchen/household/post use)	
J248		rubber sheet	
J134		other rubber pieces	
J249		rubber belts	
J125*	SUP	rubber balloons	
J126		rubber balls	
J250		rubber inner-tubes	
J251		rubber tyres	
		CLOTH/TEXTILE	
J137		clothing	
J138		shoes & sandals made of leather and/or textile	

J-CODE	SUP/FG	NAME	ITEMS COUNT
J141		cloth textile carpet & furnishing	
J140		hessian sacks/packaging	
J143		sails, canvas	
J145		other textiles	
J139		cloth textile backpacks & textile bags	

	PAPER/CARDBOARD	
J150	paper cartons/Tetrapak milk	
J151	paper cartons/Tetrapak (non-milk)	
J244	paper cups	
J245	paper food trays, food wrappers, drink containers	
J246	paper cotton bud sticks	
J247	other paper containers	
J147	paper bags	
J148	cardboard boxes	
J156	paper fragments	
J154	paper newspapers & magazines	
J158	other paper items	
J155	paper tubes and other pieces of fireworks	
J152	paper cigarette packets	
	PROCESSED/WORKED WOOD	
J159	wooden corks	
J165	wooden ice-cream sticks, chip forks, chopsticks, toothpicks	
J164	wooden fish boxes	
J163	wooden crab/lobster pots	
J162	wooden crates, boxes, baskets for packaging	
J172	other processed wooden items > 50cm	
J171	other processed wooden items 2.5 cm $\ge$ 50 cm	
J160	wooden pallets	
J167	wooden fireworks & matches	
	METAL	
J194	metal cables	
J175	metal drinks cans	
J176	metal food cans	
J181	metal tableware (e.g. plates, cups & cutlery)	
J184	metal lobster/crab pots	
J182*	metal fisheries related weights/sinkers, and lures	
J180	metal appliances (refrigerators, washers, etc.)	
J187	metal drums & barrels	
J174	metal aerosol/spray cans	
J188	other metal cans	
J190	metal paint tins	
J178*	metal bottle caps, lids & pull tabs from cans	
J195*	metal household batteries	
J177	metal foil wrappers, aluminium foil	

J199	other metal pieces > 50cm	
J198	other metal pieces 2.5cm ≥ ≤ 50cm	
J186	metal industrial scrap	
J191	wire, wire mesh, barbed wire	
J179	metal disposable BBQs	
J193	metal vehicle parts / batteries	
J130	wheels with metal hub	
	GLASS/CERAMICS	
J204	glass ceramic construction materials (bricks, tiles, cement)	
J203	glass and ceramic tableware (plates/cups/glasses)	
J207	ceramic or glass octopus pots	
J200	glass bottles	
J201	glass jars	
J208	pieces of glass/ceramic (glass or ceramic fragments ≥ 2.5 cm)	
J205	glass fluorescent light tube	
J202	glass light bulbs	
J219	other ceramic items	
J210	other glass items	
	CHEMICALS	
J216	unidentified generally dark-coloured oil-like chemicals	
J217	unidentified generally light-coloured paraffin-like chemicals	
J218	unidentified chemicals	
	FOOD WASTE	
J215	organic food waste	



This document describes the methodological approach for monitoring macrolitter on the seafloor via the use of ROV. It has been compiled based on the related methodology developed by ISPRA and the Ministry for the Environment, Land and Sea of Italy and it has been adapted within the framework of the Interreg Med PlasticBusters MPAs project to address the recent advances in the field.

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#### 9.1. Site selection

Sites should be selected to ensure that they:

- Consider areas that might accumulate litter;
- Consider areas of conservationist interest;
- Avoid areas of risk (i.e. presence of munitions).

Sites should be chosen following a two-fold approach: (i) selecting sites that meet certain criteria (e.g. are close to ports, river mouths, cities, etc.); (ii) choosing randomly from a large number of sites.

#### 9.2. Frequency and timing of surveys

At least two surveys, one in autumn and one in spring should be carried out. The proposed survey periods are:

- Autumn: October
- Spring: April

#### 9.3. Sampling unit

The sampling unit is defined by the ROV transect width and length. The surveyed area is calculated by multiplying the transect length with the visual field (width) of the ROV video. The visual field is estimated by the laser pointers scale in the video images. The estimation of litter abundance and litter interaction requires the measurement of the surveyed area.

#### 9.4. ROV operation

Given that surveys might be performed with different ROVs, with different equipment, it is important to record any ROV characteristic and instrumentation. The ROV surveys protocol should be followed as close as possible as described below:

- Transect position & orientation: Transects start at a minimum depth of 40-50 m and should be at least 200 m long. Three video transects and three replicates for each surveyed area should be performed.
- ▶ **Speed & duration:** The ROV should move along linear tracks, in a continuous recording mode, at constant low speed (< 0.3 m/s) and at a constant height from the bottom (< 1.5 m).
- Transect start and end definition: The start of the dive is defined as the moment at which the ROV dived in the seawater. The end of the dive is defined as the moment that the ROV is at sea surface / on the deck. The start of the transect is defined as the moment that the ROV is at the seafloor, and the end of the transect is defined as the moment that the ROV leaves the seafloor (off the seafloor).
- **<u>ROV characteristics</u>**: The ROV characteristics and equipment should be recorded.

#### 9.5. Litter size classes to be surveyed

The following size range classes will be reported for each recorded litter item:

- A.  $< 5 \text{ cm}^* 5 \text{ cm} = 25 \text{ cm}^2$
- B.  $< 10 \text{ cm}^{*} 10 \text{ cm} = 100 \text{ cm}^{2}$
- C. < 20cm \* 20cm = 400cm <sup>2</sup>
- D.  $< 50 \text{ cm}^{*} 50 \text{ cm} = 2500 \text{ cm}^{2}$

- E. <  $100 \text{ cm}^{*} 100 \text{ cm}^{2} = 1 \text{ m}^{2}$
- F. >  $100 \text{ cm}^{*} 100 \text{ cm} = 10000 \text{ cm}^{2} = 1 \text{ m}^{2}$

#### 9.6. Litter classification and quantification

All items observed from each video transect must be classified by type, according to the 'Joint List of Marine Litter Items Categories' prepared by the MSFD Technical Group on Marine Litter (MSFD TG ML) in close collaboration with EU Member States and the Regional Sea Conventions (Fleet et al., 2021). The manual for applying the Joint List classification system provides detailed information on how to classify litter items and a complementary photo guide helps the surveyors identify and categorise the litter items (<u>Online Photo Catalogue of the Joint List of Litter Categories</u>).

Each object observed along the transect (into the constant field of view of the camera) have to be recorded and counted, to obtain information about occurrence and abundance. All litter items observed from each video transect should be entered on the related seafloor litter monitoring sheet. Unknown litter or items that are not on the survey sheet should be noted in the appropriate "other item" box. A short description of the item should then be included on the survey sheet.

During video transect analysis, every type of litter interaction with biota and the species involved has to be noted. Interactions to recorded are just on macrofauna by visual observations only. If it is not possible to identify the organisms at species level, taxa have to be reported or at least the group. Any type of additional information has to be recorded for each item respect to interaction and impact.

The unit in which marine litter should be recorded is the number of items and it should be expressed as counts of litter items per square kilometre (litter items/km<sup>2</sup>). When it is not possible to estimate the surveyed area size (e.g. when lasers are not available), the unit in which marine litter could be expressed is items per 100 meters (items/100 m). In case of a point of litter accumulation, where it is not possible counts each single litter item, it will be identified as "litter hotspots". It will express as number of litter hotspots km<sup>-2</sup> (recommended), or number of litter hotspots km<sup>-1</sup> (mandatory).

#### 9.7. Materials and equipment

It is necessary to equip the ROV with the following items in order to carry out the ROV surveys:

- Underwater acoustic tracking position system (USBL), to provide a detailed geographical and depth position of ROV along the transects
- automatic depth system (auto depth)
- Compass
- HD Video camera (at least 1920 x 1080 pixel)
- HD Digital camera (optional)
- Laser beams at known distance, to use as a metric scale (at least two lasers).

### References

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Fleet, D., Vlachogianni, Th., Hanke, G., 2021. A Joint List of Litter Categories for Marine Macrolitter Monitoring. EUR 30348 EN, Publications Office of the European Union, Luxembourg, 2020, ISBN 978-92-76-21445-8, JRC121708. <u>https://doi.org/10.2760/127473</u>

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

Galgani et al., 2013. Guidance on Monitoring of Marine Litter in European Seas. Scientific and Technical Research series, Report EUR 26113 EN.

### Monitoring MACROLITTER on the Seafloor with ROV Data Sheet

-	Monitoring Marine Litter, Entanglement and Colonization on the Seafloor																														
Surve	Survergor Name: Location name: 0												Country:																		
Dive:					Latitud	e/longit	ude star	t:								Latitu	de/longi	tude e	nd:												
Start 1	Start Time: note:																														
					Hal	bitat		Debris	s types				Litter-f	auna inte	eraction	1		De	bris a	rea		bris sition		N of		ebris nization	s	ubstra	te		
Obs #	Time	Lat	Long	Depth (m)			n. of debris	General	ALDFG	no interaction	erage		impact-en	_		refuge	aviour	Ê	(j m j)	0 m²)				ber	Impacted species	:					
				()	Biocen osis	facies associa tion	items	waste (insert value choosing from MSFD Joint list)	(insert value choosing from MSFD Joint list)	no ini	impact -coverage	damage	mortality! necropsis	decreased mobility	other (specify)	colonization <i>I</i> refuge	adaptive behaviour	Class 1(< 1m²)	Class 2 (1 - 10 m²)	Class 3 (> 10 m²)	rolled	outstretched	species	s or colonies	epibiontic species	degree of colonizatio n		soft	soft mixed		
																											$\square$				
																	-														
_																															

Litter-fauna interaction:

- **•** *no interaction (no impact):* when there is no contact between litter and organisms
- **interaction (impact):** when there is a contact between litter and organisms:
  - coverage: when organisms and substrate portions are covered or enveloped by litter
  - entanglement: when organisms are entangled with ALDFG or other marine litter items.
  - type of damage: abrasions, wounds, broken branches (corals), epibionts, decreased mobility, mortality, etc.
  - colonization: when fouling and other sessile organisms used litter as a substrate
  - refuge: when organisms used litter as a shelter
  - adaptive behaviour: when organisms used litter as mobile shelters.

#### Litter disposition

- **rolled:** when fishing gear are all tangled up
- **outstretched**: when fishing gear is under tension (absolutely straight) or loosely lying on the seafloor with some little meandering.

#### Litter colonization/coverage:

- ▶ < 50%
- > 50%

10. Monitoring presence and impact of marine litter in biota: the Plastic Busters MPAs approach

This document describes the methodological approach for monitoring the presence and impact of Marine Litter in the Mediterranean Biodiversity (i.e., the Plastic Busters MPAs Protocols), which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML, INDICIT II Project, Barcelona Convention CORMON, and on the results of the project's testing phase.

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## THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



# **10.1.** Monitoring the presence and effects of marine litter in biota: the Plastic Busters MPAs approach

This introductive chapter describes the comprehensive methodological approach, developed within the framework of the Interreg Med Plastic Busters MPAs project, for monitoring the presence and effects of marine litter in the biodiversity inhabiting Mediterranean MPAs. This approach has been developed in order to evaluate the impact (mainly ingestion) of marine litter on both **commercially harvested** (invertebrates and fish) and **endangered species** (cetaceans, sea turtles, seabirds).

The Plastic Busters MPAs methodological approach builds on the methodologies developed by the MSFD TGML, the Barcelona Convention CORMON, the IPA-Adriatic DeFishGear project, the Interreg Med MEDSEALITTER project, the INDICIT II project, taking into account the findings of the testing phase of the Interreg Med Plastic Busters MPAs project. The resulting protocols are characterized by two main novelties:

- > The selection of a wide range of bioindicator species;
- > The development of a new diagnostic tool: the threefold monitoring approach.

#### 10.2. The marine litter impact on biodiversity: candidate biondicators selection

The selection of sentinel species to monitor the impact of marine litter on Mediterranean fauna is a crucial step for the development of harmonized sampling methods and protocols for the establishment of a consistent regional approach at Mediterranean basin scale. The selection of sentinel species, or *"candidate bioindicators"*, has to meet specific criteria and respond to the need of monitoring different habitats in Mediterranean MPAs (from coastal areas to offshore, from benthic environments to pelagic waters) at different spatial scales, and home ranges (Fossi et al. 2018).

Several sentinel species are proposed in this protocol as "candidate bioindicators" to detect the presence and impact of marine litter; they have been identified according to: 1) available data on marine litter interactions with Mediterranean marine organisms, 2) key ecological and biological criteria for selecting sentinel species, 3) key outcomes of previous projects and, 4) the main finding of the testing phase of the Plastic Busters MPAs project.

One of the main parameters considered is the marine litter (ML) **% of occurrence**: the various species analyzed are reported according to three different ranges (classes) of ML occurrence: *Low ML occurrence (0-30%), Medium ML occurrence (31-60%), High ML occurrence (61-100%)*.

This finding allows selecting species with the highest ingestion rate (reported in Table 10.1) in addition with other key ecological and biological criteria such as:

- ▶ Home range: local scale, small-scale (FAO Geographical subareas), medium-scale (Mediterranean UN Environment/MAP sub-regions) and Mediterranean Basin scale.
- Habitat: sea surface, coastal waters, open waters, seafloor.
- **Distribution** in targeted Mediterranean MPAs.
- Frequency of Occurrence of ingestion of marine litter (in bold species with % of marine litter occurrence >30%)

**Table 10-1. ML Candidate Bioindicators** proposed for each habitat, ecological compartment and home range (in **bold** species with % of marine litter occurrence >30%, \* species studied in the Plastic Busters MPAs project).

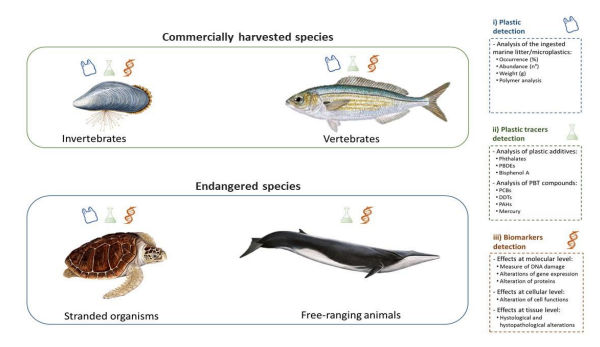
	SEA SURFACE	COASTAL WATERS	OPEN WATERS	SEAFLOOR
BASIN SCALE (Mediterranean Sea)		Puffinus yelkouan*	Balaenoptera physalus* Calonectris diomedea* Mobula mobular* Physeter macrocephalus* Thunnus thynnus Xiphias gladius	
MEDIUM-SCALE (Mediterranean UN Environment/MAP sub-regions)		Tursiops truncatus*	Caretta caretta* Chelonia mydas* Coryphaena hippurus Dermochelys coriacea Globicephala melas* Stenella -coeruleoalba* Ziphius cavirostris* Grampus griseus* Thunnus alalunga	
SMALL-SCALE (FAO GSA)	<b>Velella velella*</b> Isopods	Boops boops* Monachus monachus* Oblada melanura* Serranus cabrilla* Serranus scriba* Spicara smaris* Spondyliosoma cantharus* Trachurus trachurus*	<b>Engraulis encrasicolus*</b> <b>Sardina pilchardus*</b> Myctophium punctatum Scomber sp.	Epinephelus marginatus* Diplodus anularis* Diplodus vulgaris* Lithognathus mormyrus Merluccius merluccius* Mullus surmuletus* Pagrus pagrus* Galeus melastomus Mullus barbatus*
LOCAL SCALE		Arca noae* Modiolus barbatus* Mytilus galloprovincialis *		Holothuria forskali* Holothuria poli* Holothuria tubulosa* Arbacia lixula* Paracentrotus lividus*

In Table 10-1, the "candidate bioindicators" for each habitat and ecological compartment have been reported, also in light of the main finding from the testing phase of the Plastic Busters MPAs project. The ML candidate bioindicators are proposed according to their different frequency of occurrence, which can be used as a proxy for exposure to ML ingestion (*Medium ML occurrence (31-60%*), *High ML occurrence (61-100%*) (Fig. A1, Annex I).

#### 10.3. The threefold monitoring approach

Assessing the impact of litter on marine organisms is a challenging task. Physical and ecotoxicological effects strictly related to marine litter and, in particular, to plastics can be directly addressed in just few cases, thus calling for an integrated approach. The impact of litter on marine organisms should be assessed using a multi-tier approach, tested within the Plastic Busters MPAs project, which links marine litter ingestion detection with the physical and toxicological effects related to the ingestion of of contaminated plastic litter and the contaminants absorbed on litter and the leaching chamicals (e.g additives). The application of the threefold approach, described in the next chapters, can elucidate not only the rate of ingestion among the different bioindicators, but also the multiple sub-lethal stresses that marine litter ingestion can cause in the short and long term. Each of the three investigation tools that make up the threefold approach can be applied independently or

simultaneously to the selected candidate bioindicators. Sentinel species are subdivided into two categories: a) **commercially harvested species**; and b) **endangered species** (free-ranging and stranded marine mammals, hospitalized and stranded sea turtles) (Fig. 10.1).



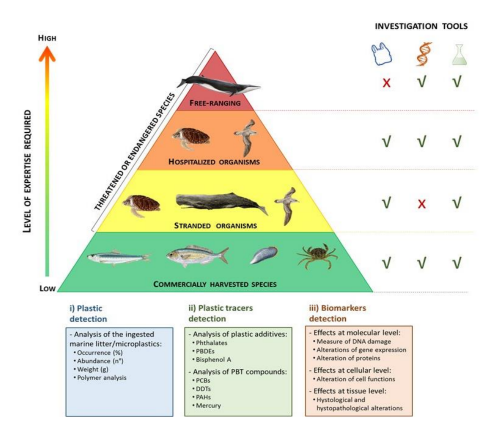
*Figure 10-1.* The threefold monitoring approach applied in the Plastic Busters MPAs project.

The implemented monitoring approach – defined as the *threefold monitoring approach* – relies on the following three types of data:

- I. analysis of the gastro-intestinal content in vertebrates/invertebrates (or of the whole organism, in the case of small invertebrates) to evaluate the marine litter ingested by the selected species, with a particular focus on plastics and microplastics. This analysis must focus on assessing the occurrence (%) of individuals that have ingested marine litter, the abundance (n° of items) of marine litter ingested per individual, the weight (g) of marine litter ingested as a total and per category of litter, the colour of litter items, as well as the polymer characterization of the plastic litter and microplastics ingested by the different individuals/species analyzed. Information on the extent to which marine biota ingests marine litter (including microplastics) is essential to determine threshold levels to define 'good environmental status' (GES) for marine litter and plastic pollution (as recommended by the EU MSFD and other regional and international regulations, i.e. Descriptor 10 of the MSFD, Ecological Objective 10 of the Barcelona Convention Ecosystem Approach).
- II. **quantitative and qualitative analyses of plastic additives** (e.g., phthalates and polybrominated diphenyl ethers-PBDEs) and Persistent, Bioaccumulative and Toxic (PBT) compounds in the tissues of bioindicators, used as "plastic tracers". The detection of plastic additives and PBT compounds that can be transferred from plastic litter to the tissues of organisms could represent the degree of accumulation of compounds related to the ingested plastic litter and the causes of its putative ecotoxicological effects. The evaluation of plastic tracers, especially in biological materials obtained in a non-lethal way in endangered species (e.g. skin biopsies), can represent a proxy of ingestion of plastic materials.
- III. analysis of the effects of marine litter and additives based on biomarker responses at different biological levels (from gene/protein expression variations to histological alterations; Omics techniques). Assessing the biological responses (alteration of a set of

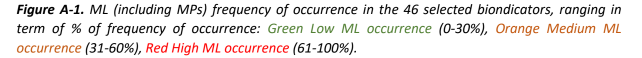
biomarkers by the measurement of endpoints) to the ingestion of marine litter and the accumulation of plastic associated compounds is crucial; this allows understanding and evaluating the extent to which marine litter and plastic ingestion pose a threat to marine organisms at individual and, ultimately, population level.

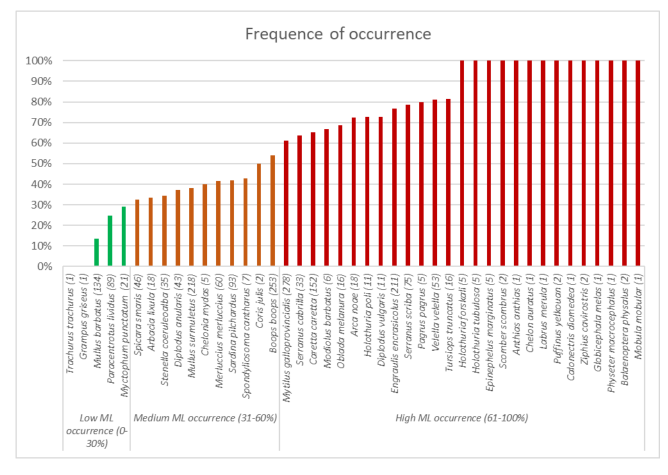
The application of the three categories of monitoring techniques (Fig. 10.2) – i) Marine litter ingested detection, ii) Plastic tracers' detection, and iii) Biomarkers detection in the candidate bioindicators– requires varying degrees of expertise, ranging from techniques easily applicable by the majority of institutions involved in marine litter monitoring (marine litter ingested detection), to the most specialized and complex ones, such as the estimation of ecotoxicological effects (plastic tracers, biomarker and Omics analysis). The gradient of expertise is described below for four typologies of organisms: a) commercially harvested species, b) stranded endangered species, c) hospitalized endangered species, d) free-ranging endangered species.



**Figure 10-2.** Level of expertise required for the detection of marine litter ingestion and impact in Mediterranean biota as adopted by the Plastic Busters MPAs project. Blue plastic bag: marine litter detection; DNA double helix: biomarker detection; green flask: contaminants (plastic tracers) detection.

#### **ANNEX I**





In order to propose a series of candidate bioindicator species to identify the presence and impact of marine litter in Mediterranean MPAs, the various species analyzed in the testing phase of the Plastic Busters MPAs project, are reported according to three different range of ML occurrence: *Green Low ML occurrence (0-30%), Orange Medium ML occurrence (31-60%), Red High ML occurrence (61-100%).* (Figure 1A). This data will subsequently be used to select the species with the highest ingestion rate which will be reported in Table 10.1 in addition with other ecological and biological parameters.



This document describes the methodological approach for monitoring the presence and effects of marine litter in invertebrates, which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML and Barcelona Convention CORMON, and on the results of the project's testing phase.

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#### 11.1. Sampling approaches

Marine invertebrate species such as filter-feeding invertebrates (e.g., mussels), and other invertebrate species (e.g., sea urchins) should be collected following any of the modalities below:

- ▶ Marine invertebrates are collected inside the study area.
- Marine invertebrates are collected in adjacent areas with similar conditions and are relocated in the study area with the use of metal cages. After a period of 3-4 weeks, they can be sampled.
- Marine invertebrates are purchased by local fishers active in the study area.

It is recommended to record the following information for each sampling site:

- **Climate variables**: Sea temperature (in °C), mean wave height, maximum wave height, mean wave period, wave direction, etc.
- Environmental variables: Sediment granulometry, nutrients, turbidity, chlorophyll-a, salinity, etc.
- Habitat Characteristics: Habitat type (e.g., sand, seagrass, algae, mats), habitat composition (% seagrass, % sand), etc.
- Coastline morphology: Beach, cliffs, estuaries, closed bay, open bay, creeks, etc.
- **Anthropogenic variables:** Anchoring allowance, diving, sewage input, fishing activities, presence of fishing gear, poaching, etc.
- **Protection status:** Protection level (fully protected, partially protected, not protected), protection status (e.g., Marine Reserve, Natural Park, Site of Community Importance), number of years before/after the establishment of protection status, etc.

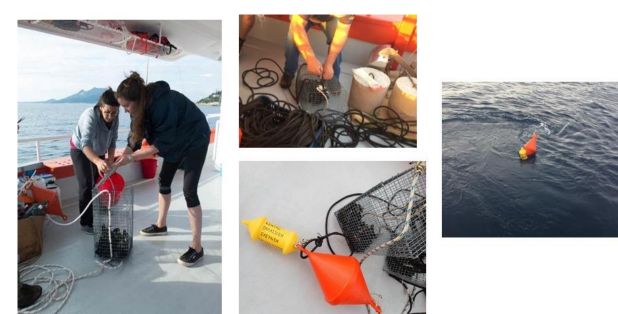


Figure 11-1. Sampling approach using translocated mussels in metal cages.

For specimens purchased from fishers the following information should be recorded: the date and time of capture, the name of the boat(s) and fishing gear used, the sampling depth. If possible, the latitude and longitude of each point where the species were captured should be recorded. If this is not possible, the area where the species were captured could be extrapolated from the Automatic Identification System (AIS).

#### 11.2. Frequency and timing of sampling

Marine invertebrates should be sampled at least once per year.

#### 11.3. Sample size

Irrespective of the chosen sampling approach (from those listed above), the minimum number of specimens sampled per sampling site should be as follows:

- Mussels: 30 specimens
- Sea urchins: 30 specimens



Figure 11-2. Sea urchins sampling.

#### 11.4. Tissues collection

To perform litter, contaminants and biomarker analyses, tissues should be removed from living organisms. Alternatively, if performing only litter and contaminant analyses, tissues can be dissected from animals frozen at -20 °C. Before the dissection of the specimens, the following information should be recorded:

- The name of the species.
- The weight of each individual (removing byssus filaments for mussels, accurate to the 4<sup>th</sup> decimal per individual).
- The length and width of each individual.
- Any visible deformations.
- The standard identification code of the animal written on the label.

Once in the laboratory, proceed either with the dissection of tissues for microplastics analysis and contaminant analysis or for biomarkers analysis or store the specimen at -20 °C or -80 °C until future dissection for microplastics or biomarkers analyses, respectively.

Dissect the following tissues (to be labelled with a unique ID for each individual):

- Hemolymph (mussels), coelomic fluid (sea urchins): hemolymph should be withdrawn from the adductor muscle of mussels using a disposable heparinized syringe with a 23G or 18G needle. The coelomic fluid should be drawn from sea urchins by a syringe inserted in the peristomal membrane around the Aristotele's lantern. Use part of the haemolymph or coelomic fluid to obtain smears and an aliquot for different biomarkers analysis (stored at 80°C).
- Digestive gland (mussels), gastrointestinal tract (sea urchins): it should be collected and weighted in aluminium paper, placed in labelled cryogenic vials, frozen in liquid nitrogen and stored at -80 °C or dry ice for biomarkers analysis. For microplastic analysis, it should be placed in aluminium foil and stored at -20 °C.
- **Gills (mussels):** they should be collected, placed in labelled cryogenic vials, frozen in liquid nitrogen and stored at -80 °C or dry ice for biomarkers analysis. For microplastic analysis, they should be placed in aluminium foil and stored at -20 °C.
- Mantle (mussels) (Fig. 11.3): it should be collected for biomarkers analyses, frozen in liquid nitrogen and stored in labelled cryogenic vials at -80 °C or dry ice. For microplastic analysis, they should be placed in aluminium foil and stored at -20 °C.
- **Gonads (mussels, sea urchins):** they should be collected for biomarker analyses, part of the tissue placed in labelled cryogenic vials, frozen in liquid nitrogen and stored at -80 °C or dry ice and another part stored in Bouin's solution.

If the dissection of the different tissues is not possible, the whole organism should be used for litter analyses.

Whole organisms should be stored at -20 °C in aluminium foil for contaminant analyses.



Figure 11-3. Dissection of mussel.

#### 11.5. Litter size classes to be surveyed

The litter size classes to be surveyed depend on the size of the investigated invertebrate. Usually for mussels and small size invertebrates, only large and small microplastics are ingested and can be detected. Litter items with their longest dimension larger than 50  $\mu$ m can be detected using the protocol described below for microplastic analyses.

#### 11.6. Litter analysis, classification and quantification

Once at the laboratory, biological samples should be first digested, then sorted and identified under a stereomicroscope with optical enhancement from 6.7x to 40.5x (Alomar et al., 2016, Nadal et al., 2016) following the protocol described in Tsangaris et al. (2021), which is the result of an intercalibration among Plastic Busters MPAs project partners.

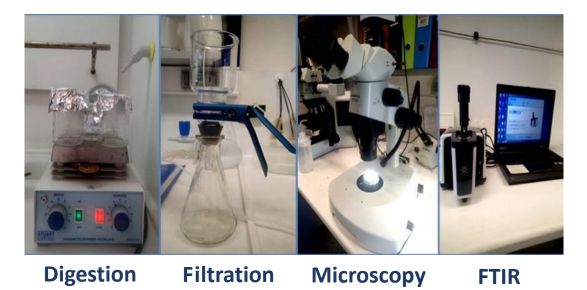
#### Microplastic analysis

- Place the tissue sample in a glass beaker; add 5 ml 10% KOH per gram of tissue wet weight (1:5 w/v).
- Cover the samples with aluminum foil and heat them up on a thermostatic water bath (50 °C) until all organic matter is removed (maximum 2 days, 12 hours heating).
- After the digestion of the organic matter, pass the samples through a metal sieve (300 μm) placed above a filtering apparatus and finally filter the sample on a fiber glass filter under vacuum (Whatman GF/C, pore size 1.2 or 1.6 μm).
- Metal sieves shall be covered with aluminium foil and filters shall be placed in aluminium foilcovered Petri dishes and dried at room temperature.
- All filtering procedures shall take place inside a laminar flow cabinet.
- Use a procedural blank sample to test for possible ambient contamination: add similar volume of 10% KOH as that used in the samples in a beaker without sample, and follow the protocol described in the steps above.
- After the digestion procedure, check the filter for plastic items with the use of a stereomicroscope.
- Photograph, count and record the type, colour and maximum length of plastic particles using an image analysis software. Categorize plastic particles according to shape, size, colour and polymer.
- Additionally, 10% of the identified items should be considered for identification using spectroscopy techniques (FT-IR, RAMAN).
- The recovery rate of microplastics by the applied extraction procedure must be tested on tissue samples enriched with specific number (e.g. 10 particles/sample) of different types of plastic particles. Use the number of particles detected after processing the sample to calculate % recovery rate of microplastics.

#### Contamination precautions

Contamination precautions are essential during all steps of the sample processing due to the ubiquitous nature of certain types of microplastics, such as synthetic fibers, that can contaminate the samples. Glass material should be used where possible and all glassware and tools (e.g. tweezers, scissors, etc.) should be rinsed thoroughly with purified water. Staff should wear natural fiber laboratory clothes. Sample processing should be done in closed areas with little ventilation and air circulation (e.g., from air conditioners). It is recommended to use covers during sample rinsing and filtration (e.g., glove bag, laminar flow cabinet or other closed cover) and to cover filters with glass lids during observation under the stereomicroscope. Procedural blank samples should be used throughout the entire sample processing. During the analysis procedure, two glass petri dishes should be placed at each side of the stereomicroscope and

checked for microplastics before and after each sample. A 100% cotton laboratory coat shall be worn at all times during the procedure.



*Figure 11-4.* Main steps of invertebrate sample processing for microplastic detection (from Tsangaris et al. 2015).

#### Collection of data

For each species an assessment is made of:

- 1. Frequency of occurrence (%) of ingested microplastics, calculated as the percentage of the individuals examined with ingested microplastics.
- 2. Abundance (N) of macro and microplastics ingested per individual (average number of items/individual) for each species, calculated as a total and per category. Given currently existing inconsistencies in the literature regarding reporting the abundance of ingested litter, it is recommended to report average number of items per individual, considering both all the individuals examined and those solely found with ingested litter.

For each individual organism, the following information on ingested litter shall be reported:

- 1. The number, length, weight and nature of the polymer (10%) of the items examined.
- 2. The recovery rate of microplastics.

#### **11.7.** Analysis of plastic tracers and PBTs

#### Plastic additives

The compounds to be detected are:

- <u>*Phthalates:*</u> a group of chemicals widely used (e.g., plastic additives) to make plastics more flexible and harder to break; they can interfere with the endocrine system (Baini et al., 2017).
- <u>Bisphenol A</u>: used in the production of polycarbonate, can have endocrine disrupting effects (Crain et al., 2007; Halden, 2010; Oehlmann et al., 2009) and the styrene and polyvinyl chloride monomer, used in the production of polystyrene and polyvinyl chloride (PVC), can be carcinogenic and/or mutagenic (Lithner et al., 2011; Papaleo et al., 2011; Xu et al., 2004).

• <u>Polybrominated diphenyl ethers:</u> they belong to the group of brominated flame retardants (BFRs), which are used in various polymeric materials (e.g., plastic parts, resins, textiles, and other substrates) to reduce their fire hazards (BSEF 2003; Król et al. 2012).

#### Persistent, bioaccumulative and toxic substances (PBTs)

In addition to the plastic additives that may leach from the plastic items released into the marine environment, persistent bioaccumulative and toxic substances (PBTs) (e.g. organochlorine compounds OCs, PAHs and PBDEs) and metals (e.g., lead, copper and cadmium) that are present in the seawater tend to accumulate in the surface of plastic items.

Depending on the compounds and the tissues to be analysed, different methods should be applied to detect the presence of plastic-related contaminants in invertebrates (Annex II, Table A.1, Fig. 11.5).

#### 11.8. Biomarkers analysis

The toxicological effects associated with the presence of marine litter can be evaluated using a set of diagnostic and prognostic methodologies, by means of biomarkers. A non-exhaustive list of existing biomarker approaches and plastic tracers' contaminants that are usually applied in invertebrates analyses is reported in Annex II (Table A.2) and Figure 11.5.

Biomarkers have been selected on the basis of the level of biological responses and in relation to the main effects related to marine litter/microplastics ingestion. The selected biomarkers can diagnose different impacts related to: a) physical damages/effects of marine litter, b) exposure to/effect of chemical tracers, and c) exposure to/effect of adsorbed chemicals.

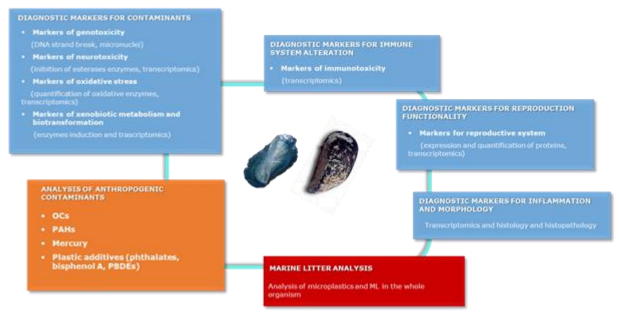
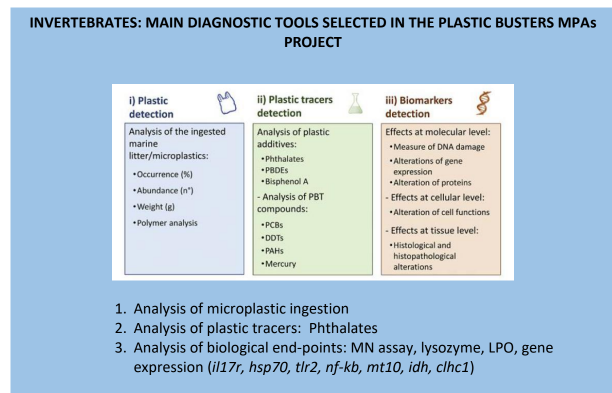


Figure 11-5. A three-fold approach to detect the marine litter presence and impacts to invertebrates.

Starting from this initial list and building on the findings of the testing phase of the Plastic Busters MPAs project, the most suitable diagnostic tools to detect the presence and impact of ML on invertebrates are proposed here below.

**Table 11-1.** Main diagnostic tools selected in the Plastic Busters MPAs project to detect the presence and impact of ML in invertebrates.



#### 11.9. Materials & Equipment

#### Material for sampling

- Camera
- Containers for samples, zipped bags, cool boxes
- Garbage bag
- Gloves
- Disposable scalpels, fine forceps, scissors and tweezers for dissection
- Dissection board
- Measuring decimetre
- Pen/pencil/ Permanent marker
- Sampling sheets
- Aluminium foil
- Cryoboxes
- Cryovials
- Liquid nitrogen Dewar (in alternate dry ice)
- Paper and block-notes
- Paper towels

#### Material for microplastic analysis

- Distilled water
- Permanent marker
- 10% KOH
- Disposable scalpels, fine forceps, scissors and tweezers for dissection
- Precision tweezers (fine and pointed) for micro-plastic handling on filters and FTIR

- Glass petri dishes
- GFC filters 0.2, 1.2 or 1.6 μm 47 mm diameter for filtration
- Aluminum foil
- 150 and 250 ml glass beakers
- 250 ml conical flasks
- 100 ml measuring glass cylinder
- Magnetic stirrer
- Hot plate
- Glove bag, laminar flow cabinet or other closed cover
- Vacuum filtration system with ramp
- Analytical balance
- Stereomicroscope with image analysis software
- FTIR or Raman spectroscopy with associated analysis software

#### References

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

Guidance on Monitoring of Marine Litter in European Seas, 2022. A guidance document within the Common Implementation Strategy for the marine Strategy Framework Directive - Marine Litter Impact on Biota. MSFD Technical Group on Marine Litter - draft.

## Monitoring Microlitter in biota: mussels

Sampling date and time	Sampling site	GSA	Sampling method	Depth	Coordinates	
					Latitude	Longitude

ID code Species	Shell length Sex (cm)	Shell length (cm)	Shell weight	Soft tissue	Digestive gland weight (g)	Gills	Hemolymph		Mantle	
ib code	species	Species Sex	(ciii)	(g)	weight (g)	weight (g)	GIII3	MN test	(ul)	Mantie

## Monitoring Microlitter in biota: sea urchins

Sampling date and time	Sampling site	GSA	Sampling method	Depth	Coordinates	
					Latitude	Longitude

ID code	Species	Sex	Weight (g)	Diameter (cm)	Soft tissues weight (g)

Notes and remarks:

#### **ANNEX II**

	CHEMICAL COMPOUND	TISSUE/SAMPLE	ANALYTICAL METHOD
	Phthalates	Muscle, whole organism	Baini et al., (2017), Fossi et al., (2016), Savoca et al., (2018), Avisar et al., (2019), Lo Brutto et al., (2021),
PLASTIC ADDITIVES	Bisphenol A	Muscle, whole organism	Ballesteros-Gómez et al., (2009), Lo Brutto et al., (2021)
	Polybrominated diphenyl ethers	Muscle, whole organism	Muñoz-Arnanz et al., (2016), Cruz et al., (2019), Cruz et al., (2020)
	Polycyclic aromatic hydrocarbons	Muscle, whole organism	Marsili et al., (2001), León et al., (2013), Benedetti et al., (2014)
ADSORBED CONTAMINANTS	Organochlorine contaminants	Muscle, whole organism	Marsili and Focardi, (1997), León et al., (2021)
	Mercury	Whole organism, muscle	Correa et al., (2013), Fattorini et al., (2008), Besada et al., (2011), León et al., (2021)

 Table A-1. Tissues and methods to be used to detect plastic tracers in invertebrates.

EFFECT	TISSUE	TEST		
GENOTOXICITY	Hemolymph, digestive gland	Comet assay (Revel et al., 2019) (*) Mn test (Avio et al., 2015) (*)		
OXIDATIVE STRESS	Digestive gland	LPO, CAT, SOD, GST, GSH, GR, GPX (Revel et al., 2019) (*) qPCR GPX, SOD, CAT (Ravel et al., 2019)		
IMMUNOTOXICITY	Gills, Mantle, Digestive gland	CASP, TRAF, Transcriptomics (Avio et al., 2015; Revel et al., 2019) (*) Transcriptomics (Gardon et al., 2020) qPCR LYS, CASP3 (Paul- Pont et al, 2016)		
REPRODUCTION	Gonads	Gamete Quality and Larval Development (Sussarellu et al., 2016) (*)		
HISTOPATHOLOGY INFLAMMATION AND MORPHOLOGY	Digestive gland	Histopathology, histology (Avio et al., 2015) (*)		
XENOBIOTIC METABOLISM AND BIOTRANSFORMATION	Digestive gland, whole organism	Porphyrins (Grandchamp et al. 1980; Guerranti et al. 2014) (*) EROD (Zhang et al., 2019) Transcriptomics (Gardon et al., 2020) (*)		
NEUROTOXICITY	Whole organisms, muscle, gills	AChE activity (Magni et al., 2018) (*)		
CELLULAR STRESS	Whole organisms, muscle, hemolymph, digestive gland	Lysosomal membrane stability-LMS (Canesi et al 2015) (*) IDH (Oliveira et al., 2013) (*) Transcriptomics (Détréé et al. 2018) qPCR IDH, HSP70 Détréé et al. 2017)		

 Table A-2. Effects measured in invertebrates by the biomarker approach.

(\*) effects detected after laboratory or field exposure with MPs or plastic-related contaminants.



This document describes the methodological approach for monitoring the presence and effects of marine litter in fish, which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML, INDICIT II Project and Barcelona Convention CORMON, and on the results of the project's testing phase.

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#### 12.1. Sampling approaches

Fish species should be sampled following one of the following approaches depending on the type of analysis to be performed:

- ▶ For the analysis of litter and associated contaminants, fish species (dead) can be purchased by local fishers active in the study area.
- ▶ For the analysis of litter, associated contaminants and biomarkers, fish species (live) should be collected in the study area via a dedicated sampling campaign.

It is recommended to record the following information for each sampling site:

- **Climate variables**: Sea temperature (in °C), mean wave height, maximum wave height, mean wave period, wave direction, etc.
- Environmental variables: Sediment granulometry, nutrients, turbidity, chlorophyll-a, salinity, etc.
- Habitat Characteristics: Habitat type (e.g., sand, seagrass, algae, mats), habitat composition (% seagrass, % sand), etc.
- Coastline morphology: Beach, cliffs, estuaries, closed bay, open bay, creeks, etc.
- **Anthropogenic variables:** Anchoring allowance, diving, sewage input, fishing activities, presence of fishing gear, poaching, etc.
- **Protection status:** Protection level (fully protected, partially protected, not protected), protection status (e.g., Marine Reserve, Natural Park, Site of Community Importance), number of years before/after the establishment of protection status, etc.

#### 12.2. Frequency and timing of surveys

Frequency of sampling is at least once per year, taking into account seasonality.

#### 12.3. Sample size

A minimum of 30 individuals per fish species should be sampled at each site, preferably for each environmental compartment (i.e., benthic, demersal, pelagic). Specimens of endangered species (e.g. Manta ray) occasionally found stranded can also be analyzed in very small numbers.

#### 12.4. Tissues collection

To perform litter, contaminants and biomarker analyses, tissues should be removed from living organisms. Alternatively, if performing only litter and contaminant analysis, tissues can be removed from animals frozen at -20 °C. Before the dissection, the following information should be recorded for each fish sample:

- Date and time of capture
- Name of sampling location
- Name of the boat(s) providing the samples
- Sampling gear
- Latitude and longitude of each point where species are captured
- Sampling depth
- Sample size: number of individuals sampled.

Immediately after sampling, rinse the fish and label the fish samples with a unique ID for each individual.

Before the dissection of the fish:

- Record the name of the species
- Weigh the whole fish
- Measure the total length of the fish
- Record any visible deformations
- Record the gender (if possible)
- Record the maturity stage

To avoid airborne contamination, it is recommended to dissect the specimen and take tissue samples in the laboratory, under controlled conditions.

#### Dead organisms

Before dissection, thaw fish in the laboratory (if previously stored at -20 °C) at room temperature.

Collect the following tissues:

- Gastrointestinal tract (GI) for litter analysis: whole GI in aluminium foil at -20 °C.
- Muscle for contaminants analysis: about 1g in aluminium foil stored at -20 °C.
- Liver for contaminant analysis: about 1g in aluminium foil stored at -20 °C. Weight the liver for somatic liver index (SLI) evaluation.

Each tissue stored in aluminium foil must be labelled with a unique ID for each individual.

#### Live organisms

Keep the sampled live animals on board, in seawater with oxygenators, transport and dissect the animals in the laboratory. Alternatively, animals can be dissected on board. Before dissection, anaesthetise the animals following related guidelines.

Extract the following tissues:

- Blood samples for biomarker analysis: the blood should be withdrawn from the caudal vein using a disposable heparinized syringe. Use part of the blood to obtain blood smears and centrifuge an aliquot of the blood to obtain plasma samples.
- Liver for biomarker analysis: about 1g in aluminum paper, weight the liver for somatic liver index (SLI) evaluation, freeze in liquid nitrogen or dry ice in cryovials and store at -80°C.
- Kidney for biomarker analysis: frozen in liquid nitrogen or dry ice in cryovials and stored at 80°C.
- Gills for biomarker analysis: frozen in liquid nitrogen in cryovials or dry ice and stored at 80°C.
- Muscle sample: an aliquot frozen in liquid nitrogen or dry ice in cryovials and stored at -80 °C for biomarker analysis, and a part of the sample for contaminants analysis in aluminium foil, stored at -20 °C.
- Gastrointestinal tract (GI): whole GI in aluminum foil stored at -20°C for litter analysis.

Each tissue stored in aluminium foil or cryovial/eppendorf must be labelled with a unique ID for each animal.



Figure 12-1. Fish blood sampling (live fish).

#### 12.5. Litter size classes to be surveyed

Litter items smaller than 5 mm can be classified in different size classes, large and small microplastics. Lowest limit for microplastic should be 100  $\mu$ m. For large fish species (e.g. swordfish, tuna), where is possible, under notes in datasheets, the items should be described and assigned a litter category number using the "Joint List" developed by the MSFD TGML group (Fleet et al., 2021).

#### 12.6. Litter analysis and classification

- Place GI (stomach and intestine) in a glass petri dish or beaker. Be careful to annotate the fish ID in each petri-dish/beaker.
- Weigh and rinse the GI with purified water (e.g. milli Q).
- Place a filter paper in a petri dish (blank sample) in the working area during fish dissection to test airborne contamination.

#### Macrolitter detection

- For macrolitter and gut content analysis, cut open the stomach and intestine, remove stomach and intestine contents and weigh separately.
- Sort prey or litter items into separate categories under a stereomicroscope, taking care of recording their weight.
- Measure the size of litter items and classify litter categories classified according to the JointList of Litter Categories of the MSFD Technical Group on Marine Litter.

In addition, the following parameters should be recorded:

 Record for all categories (litter and other elements) the dry mass (grams, precision 0.01 g) of each category: dry the sample at room temperature for 24 h minimum or in a stove at 35 °C for 12 h.

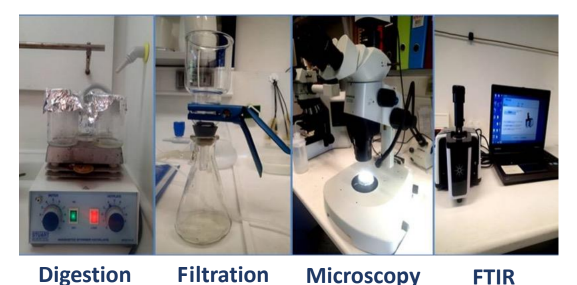
# Microlitter detection

For microlitter analysis apply the following digestion procedure described in Tsangaris et al. (2021), which is the result of an intercalibration among the PB-MPAs project partners. To avoid losing content, digest the entire GI and not just its content. The GI can be divided in two subsamples for faster digestion since time required for digestion depends on the amount of tissue to be digested.

# Microplastic and macroplastic analysis

For microplastic and microplastic analyses apply the following procedure:

- Place the entire GI in a glass beaker, add 5ml 10% KOH per gram of tissue wet weight (1:5 w/v).
- Cover the samples with aluminum foil and heate on thermostatic water bath (50 °C) until all organic matter is removed (maximum 2 days, 12 hours heating).
- After the digestion of the organic matter, pass the samples through a metal sieve (300 μm) placed above a filtering apparatus and finally filter under vacuum onto a fiberglass filter (Whatman GF/C, pore size 1.2 or 1.6 μm).
- Metal sieves should be covered with aluminum foil and filters must be placed in aluminum foil-covered Petri dishes and dried at room temperature.
- Use a procedural blank sample to test for possible ambient contamination: add similar volume of 10% KOH as that used in the samples in a beaker without sample, and follow the protocol described in the steps above.
- After the digestion procedure, check the filter for plastic items with the use of a stereomicroscope.
- Photograph, count and record the type, colour and maximum length of plastic particles using an image analysis software. Categorize plastic particles according to shape, size, colour and polymer.
- The recovery rate of microplastics by the applied extraction procedure must be tested on tissue samples enriched with specific number (e.g. 10 particles/sample) of different types of plastic particles. The number of particles detected after sample processing is used to calculate % recovery rate of microplastics.
- To avoid contamination, carry out filtration under a cover (e.g. glove bag, laminar flow cabinet or other closed cover).
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.
- Test recovery of microplastics by the applied extraction procedure on fish tissue samples enriched with specific number (e.g. 10 particles/sample) of different plastic particles of known polymer type and size (positive controls, minimum number: The number of particles detected after the processing of these samples as described above, should be used to calculate % recovery of microplastics.



*Figure 12-2.* Main steps of fish sample processing for microplastic detection (from Tsangaris et al. 2015).

#### Contamination precautions

Contamination precautions are essential during all steps of the sample processing due to the ubiquitous nature of certain types of microplastics, such as synthetic fibers, that can contaminate the samples. Glass material should be used where possible and all glassware and tools (e.g. tweezers, scissors, etc.) should be rinsed thoroughly with purified water. Staff should wear natural fiber laboratory clothes. Sample processing should be done in closed areas with little ventilation and air circulation for example from air conditioners. Samples should be covered by foil paper during digestion and when not in use. It is recommended to use covers during sample rinsing and filtration (e.g. glove bag, laminar flow cabinet or other closed cover) and during all steps of samples processing (e.g. dissection, examination under the stereomicroscope). Procedural blank samples should be throughout the entire sample processing. During the analysis's procedure, two glass petri dishes should be placed at each side of the stereomicroscope and checked for microplastics before and after each sample. A 100% cotton laboratory coat shall be worn at all times during the procedure.



Figure 12-3. Filtration of digested sample in a glove box.

# Collection of data

For each organism an assessment is made of the:

- 3. Frequency of occurrence (%) of ingested macro- and microplastics for each species is calculated as the percentage of the individuals examined with ingested microplastics.
- 4. Abundance (N) of macro- and micro-plastics ingested per individual (average number of items/individual) for each species is calculated as a total and per category. Since currently there are inconsistencies in the literature in reporting abundance of ingested litter, it is recommended to report average number of items per individual both considering all individuals examined and only individuals found with ingested macrolitter and microlitter.
- 5. The percentage of the individuals affected in relation with the individuals of the whole sample examined (all species).

For each organism data on litter ingested is reported:

- 1. Characteristics of the litter found (colour, shape, size and polymer) in each specimen according to the "MSFD Protocol for the monitoring of microliter ingested by marine fish".
- 2. The number, length, weight and nature of the polymer (10%) of the items examined for each species.
- 3. Recovery rate of microplastics.

# 12.7. Analysis of plastic tracers and PBTs

# Plastic additives

The compounds to be detected in different tissues/fluid are:

- <u>Phthalates:</u> a group of chemicals widely used as additives to make plastics more flexible and harder to break; they can interfere with endocrine system (Baini et al., 2017).
- <u>Bisphenol A</u>: used in the production of polycarbonate, can have endocrine disrupting effects (Crain et al., 2007; Halden, 2010; Oehlmann et al., 2009) and the styrene and polyvinyl chloride monomer, used in the production of polystyrene and polyvinyl chloride (PVC), can be carcinogenic and/or mutagenic (Lithner et al., 2011; Papaleo et al., 2011; Xu et al., 2004).
- <u>Polybrominated diphenyl ethers</u>: they belong to the group of brominated flame retardants (BFRs), which are used in various polymeric materials such as plastic parts, resins, textiles, and other substrates to reduce their fire hazards (BSEF 2003; Król et al. 2012).

# Persistent, bioaccumulative and toxic substances (PBTs)

In addition to the plastic additives that may leach from plastics when released into the marine environment, plastics tend also to adsorb in their surface persistent bioaccumulative and toxic substances (PBTs) (e.g. organochlorine compounds OCs, PAHs and PBDEs) and metals (e.g., lead, copper and cadmium) that are present in the seawater.

Depending on the compounds and the tissue to be analysed, different methods should be applied to detect the presence of plastic-related contaminants in the fish species (Annex III, Table A.3, Fig. 12.4).

# **12.8.** Biomarkers analysis

The toxicological effects associated with the presence of marine litter can be evaluated using a set of diagnostic and prognostic methodologies, by means of biomarkers. A non-exhaustive list of existing biomarker approaches and plastic tracers' contaminants that are usually applied in fish analyses is reported in Annex III (Table A.4).

Biomarkers have been selected on the basis of the level of biological responses and in relation to the main effects related to marine litter/microplastics ingestion. The selected biomarkers can diagnose different impacts related to: a) physical damages/effects of marine litter, b) exposure to/effect of chemical tracers, and c) exposure to/effect of adsorbed chemicals.

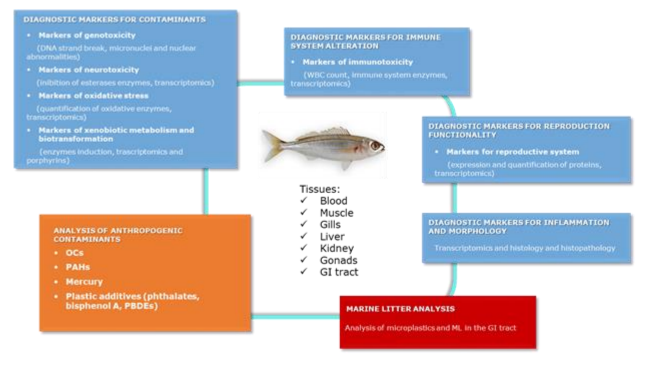
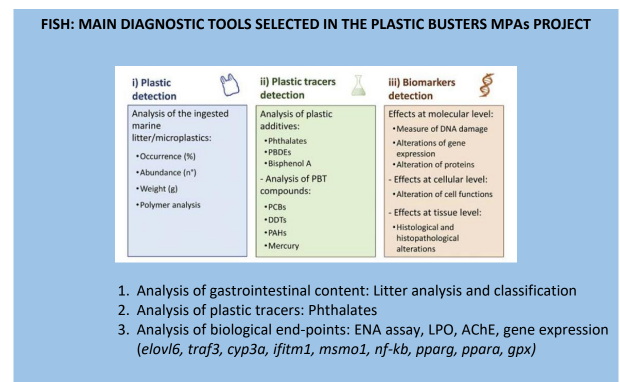


Figure 12-4. A three-fold approach to detect the marine litter presence and impacts to fish species.

**Table 12-1.** Main diagnostic tools selected in the Plastic Busters MPAs project to detect the presence and impact of ML in fish.



Starting from this initial list and building on the findings of the testing phase of the Plastic Busters MPAs project, the most suitable diagnostic tools to detect the presence and impact of marine litter on fish are proposed here below.

# 12.9. Materials & Equipment

# Material for sampling

- Camera
- Containers for samples, zipped bags, cool boxes
- Garbage bag
- Gloves
- Disposable scalpels, fine forceps, scissors and tweezers for dissection
- Dissection board
- Measuring decimetre
- Pen/pencil/ Permanent marker
- Sampling sheets
- Aluminium foil
- Cryoboxes
- Cryovials
- Liquid nitrogen Dewar (in alternate dry ice)
- Paper and block-notes
- Paper towels

# Material for macroplastics and microplastics analysis

- Distilled water
- Permanent marker
- 10% KOH
- Disposable scalpels, fine forceps, scissors and tweezers for dissection
- Precision tweezers (fine and pointed) for micro-plastic handling on filters and FTIR
- Petri dishes
- GFC filters 0.2, 1.2 or 1.6 μm 47 mm diameter for filtration
- Aluminum foil
- 150 and 250 ml glass beakers
- 100 ml measuring glass cylinder
- Magnetic stirrer
- Hot plate
- Glove bag, laminar flow cabinet or other closed cover
- Vacuum filtration system
- Analytical balance
- Stereo microscope with image analysis software
- FTIR or Raman spectroscopy with associated analysis software

# References

Guidance on Monitoring of Marine Litter in European Seas, 2022. A guidance document within the Common Implementation Strategy for the marine Strategy Framework Directive - Marine Litter Impact on Biota. MSFD Technical Group on Marine Litter - draft.

INDICIT II, Final report, 2021. Deliverable D1.6, of the European project "Implementation of the indicator of marine litter on sea turtles and biota in Regional Sea conventions and Marine Strategy Framework Directive areas".

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

# Monitoring Marine Litter (Macro-Micro) in biota: dead fish

Sampling date and time	Sampling site	Boat name	GSA	Sampling gear	Depth	Coordinates	
						Latitude	Longitude

ID code	Species	Sex	Total length (cm)	Total weight (g)	GI weight (g)	Muscle	Liver weight (g)

Notes and remarks:

# Monitoring Marine Litter (Macro-Micro) in biota: live fish

Sampling	g date and tir	ne	Sampling loca	tion	Вс	oat name	GSA	Sam	oling gear	Depth			С	oordinates		
												Lati	tude		Longitud	е
ID code	Species	Sex	Total length (cm)	Fo length		Total weight (g)	GI weight (g)	Muscle	Liver weight (g)	N°. of liver aliq.	Bile	Brain	Kidney	Gonad weight (g)	Blood smears	Plasma

Notes and remarks:

Rack (N <sub>2</sub> )						

# **ANNEX III**

	CHEMICAL COMPOUND	TISSUE/SAMPLE	ANALYTICAL METHOD	
	Phthalates	muscle, liver, whole organism	Baini et al., (2017), Fossi et al., (2016), Savoca et al., (2018)	
		Blood	Takatori et al., (2004)	
PLASTIC ADDITIVES		Muscle	Ballesteros-Gómez et al., (2009), Barboza et al., (2020)	
	Bisphenol A	Blood	Cobellis et al., (2009)	
	Polybrominated diphenyl ethers	Muscle, liver, blood	Muñoz-Arnanz et al., (2016), Bartalini et al., (2019), Ameur et al., (2020), Corsolini et al., (2008)	
	Polycyclic aromatic hydrocarbons	Muscle, liver, blood	Marsili et al., (2001), Frapiccini et al., (2020)	
ADSORBED CONTAMINANTS	Organochlorine contaminants	Muscle, liver, blood	Marsili and Focardi, (1997), Bartalini et al., (2019), Garcia-Garin et al., (2020)	
	Mercury	Blood, muscle	Correa et al., (2013), Barboza et al., (2018)	

 Table A-3. Tissues and methods to be used to detect plastic tracers in fish.

EFFECT	TISSUE	TEST
GENOTOXICITY	Blood	Comet assay (Molino et al., 2019) (*) Mn test (Bolognesi et al., 2006) ENA assay (Pedà et al., 2022); (Pacheco and Santos, 1997)
	Liver, kidney, gill	CAT, GST, LPO, GPX, GR, GSH (Pedà et al., 2022 ; Yu et al., 2018) (*) qPCR NRF2, CAT, SOD (Espinosa et al., 2019)
OXIDATIVE STRESS	Plasma	LPO (Pedà et al., 2022 ; Campani et al., 2020, Casini et al., 2018) CAT (Pedà et al., 2022)
IMMUNOTOXICITY	Blood, liver	Total and differential white blood cells (WBC) count (Casal and Orós, 2007; Davis et al., 2008; Caliani et al., 2019) H:L ratio (Caliani et al., 2019) Respiratory burst (Secombes, 1990; Caliani et al., 2019) TAS assay (Miller et al., 1993; Caliani et al., 2019) Lisozyme enzyme (Keller et al., 2006; Caliani et al., 2019) (*) Transcriptomics (Limonta et al., 2019) qPCR IL1B, IL8 CASP3 (Espinosa et al., 2019); CASP8, CASP9, TRAF (Karami et al. 2017)
REPRODUCTION	Plasma, Gonads, Liver	CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015) (*) Vitellogenin (Fossi et al., 2004), Vtg (Mak et al., 2019)
	Plasma	Vitellogenin (Herbst et al., 2003) CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*)
HISTOPATHOLOGY INFLAMMATION AND MORPHOLOGY	Liver, kidney, gill	Histopathology,histology (Pedà et al. 2016; Karami et al. 2017; Batel et al., 2018) (*)
XENOBIOTIC METABOLISM AND	Liver, blood, bile	Porphyrins (Grandchamp et al. 1980; Guerranti et al. 2014) (*) Bile metabolites (Oliveira et al 2013) (*) EROD (Zhang et al., 2019) (*)
BIOTRANSFORMATION	Blood, skin, liver	CYP1A; AHR, CYP3A (Fossi et al. 2014, Panti et al. 2011; Rochman et al., 2013) (*) Porphyrins (Guerranti et al., 2014) (*)
NEUROTOXICITY	Brain, muscle, plasma	AChE, BChE (Barboza et al., 2018) (*)
	Whole organisms, muscle	Lysosomal membrane stability-LMS (Canesi et al 2015) (*) IDH (Oliveira et al., 2013) (*)
CELLULAR STRESS	Blood, skin, liver, kidney	PPARA, PPARG, HSP70, GPX, E2F1 (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*) Gamma glutamyl transferase (GGT) (Nematdoost Haghi and Banaee, 2017) (*) Cortisol and corticosterone (Flower et al., 2015) LDH (Nematdoost Haghi and Banaee, 2017) (*)

(\*) effects detected after laboratory or field exposure with MPs or Plastic-related contaminants.



This document describes the methodological approach for monitoring the presence and effects of marine litter in sea turtles, which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML, INDICIT II Project and Barcelona Convention CORMON, and on the results of the project's testing phase.

PREPARED BY

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



### 13.1. Species sampling

The presence and impact of marine litter in Mediterranean sea turtles (*Caretta caretta, Chelonia mydas* and *Dermochelys coriacea*) can be investigated and/or monitored in:

- Dead organisms that may have been stranded ashore, captured by fishers (by-catch), found at sea or died at a recovery centre.
- Living organisms that have been hospitalized in a rescue centre.

### 13.2. Description of investigated turtle and biometric measurements

#### Description of the investigated turtle

Identify the species of the observed marine turtle and in case of doubt about the species identification, refer to an identification guide (e.g. <u>www.cites.org</u>). If the species can't be identified, note it as non-identified (NI) on the observation sheet.

The sea turtles are protected species, therefore only authorized people can handle live and dead animals or parts of them. Upon finding the animal, its management and recovery should be reported and coordinated with the responsible National Authorities. Note that a CITES permit is asked if a specimen or sample has to be sent/received or to be transported.

Indicate the presence and code number of tag, if present, otherwise, note "NO".

Note the date of discovery (dd/mm/yyyy), the location of discovery and the coordinates if available (X, Y: in decimal degrees, or specify the coordinate system); the name and contacts (phone, mail) and institution of the person in charge of the recovery shall also be noted.

Take a photo of the animal before handling.

All the sea turtle's data should be noted down in a sampling sheet (see paragraph 12.9).

#### **Biometric Measurements**

Morphometric measurements should be collected before the necropsy (if the animal is dead) or tissue collection (for specimens recovered in a rescue centre). Standard Curved Carapace Length (CCL), notch to tip is mandatory, while other measurements are optional (in centimetres, precision 0.01 cm) in addition to this measures weight (kilograms, precision 0.01 g) and sex of the specimen should be recorded:

Biometric me	asureme	ent	
1. CCLst	cm	6. HW	cm
2. CCWst	cm	7. PTL	cm
3. SCLst	cm	8. CTL	cm
4. SCWst	cm	9. CaCL	cm
5. CPL	cm		
Weight (kg)			
Sex			
JEX			

**Figure 13-1.** Biometric parameters to be measured mandatory (Standard Curved Carapace Length – CCLst, weight and sex) and optional (Standard Curved Carapace Width – CCWst; Standard Straight Carapace Length – SCLst; Straight Carapace Width – SCWst; Curved Plastron Length – CPL; Head Width – HW; Plastron Tail Length – PTL; Cloaca Tail Length – CTL; Carapace Cloaca Lenght - CaCL) (from INDICIT, 2018).

#### 13.3. Conservation/health status of the organism

- With regards to the status of the organism, two cases are possible: the turtle may be live, or dead. In the case of live animals (Level 1) biological samples (blood, carapace, plasma, biopsy, faeces) can be collected for biomarker and chemical analyses. In case of dead animals, 4 different situations can be observed: in animals that have just died (< 2 hours post mortem), gastrointestinal (GI) system is adequate for litter ingestion analysis, and other tissues (muscle, liver) can be used for biomarker and chemical analyses.</li>
- Levels 2 (fresh), 3 (partial decomposed) and 4 (advanced decomposed): are adequate for litter ingestion analysis (in GI) from necropsies and chemical analysis.
- Level 5 (mummified): the litter ingestion analysis is not possible because the individuals have usually lost their gastro-intestinal material.







Level 1- Alive Lev

Level 2- Fresh Dead recently, turtle in good condition Level 3- Partial Internal organs still in good conditions. Autolysis. Bad smell. Colour changes in skin.





Level 4- Advanced

Level 4- Mummified

Figure 13-2. Conservation status codes for stranded organisms (from UNEP/MAP, 2019).

# Discovery circumstances

Note the circumstances among the different categories:

- Stranded: animal found stranded on the beach or in the shoreline.
- By-catch/fisheries: animal accidentally captured by fishers.
- At sea: dead animal found on sea surface.
- Dead RC: the animal arrived live but died during its hospitalization in the rescue centre.
- Unknown = unknown

# Possible cause of morbidity and mortality, type of impact

If possible, the type of interaction with human activities and impact observed or suspected on dead or live stranded individuals should be deduced from external or organs observations during the necropsy and complemented with veterinarian examinations.

Also, an inspection of the oral cavity should be conducted for the presence of foreign material. Then one of the following 9 categories should be selected and noted; the remarks box should be completed with the help of the pathologist (if this is requested):

- 1. Bycatch/Fisheries related: ingested hook, decompression sickness, individual trapped in a fishing gear, individual drowned in a fishing gear;
- 2. Entanglement in debris: entanglement in litter other than related to fishing activity. Please fill the column "Entanglement type" and "Litter causing entanglement";
- 3. Ingestion of litter: digestive obstruction or occlusion, perforation or other impacts;
- 4. Anthropogenic trauma: Collision with a boat or a propeller, individual beaten with knife, stick or harpoon, poaching;

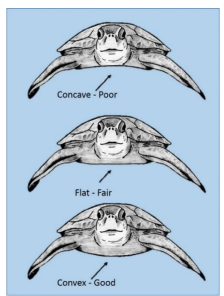
- 5. Natural predation: usually shark attack;
- 6. Natural disease: buoyancy trouble, cachexia, dermatitis, conjunctivitis, rhinitis;
- 7. Oils: Ingestion or external impregnation with oils;
- 8. Unidentified: Impossible to know the cause of death/stranding, no remarkable damages, injury or disease.
- 9. Other: Please specify in the column "Notes".

# Main injuries

In case of injuries, the *main type of injury* (bone fracture, amputation, slightly or deep cuts, throttle, abrasion or other) and the *affected body part* should be reported. If the individual has been found entangled in litter, the type of material in which the sea turtle was found should be specified, according to the following categories: *Pieces of net (N), Monofilament line (nylon) (L), Rope or pile of ropes (R), Plastic bag (Pb), Raffia (Rf), Other plastics (Ot), Multiple materials (Mu), Unknown (Unk).* 

# Health status of live animals

Note the health status according to the level of body condition by visual observation of the plastron shape.



**Figure 13-3.** Health status from visual observation of plastron shape: concave plastron poor health status; flat plastron fair health status; convex plastron good health status (from Thomson et al., 2009).

# Other descriptive parameters

Visual inspection of the animal's fat reserves at the neck is recommended. For dead individuals, this can be verified when opening the plastron according to the quantity of fat recovering the abdominal muscles. Choose among the 3 categories:

- Thin (sunken neck)
- Fat
- Normal

If possible, the sex (male or female) should be noted, which can be determined by gonads analysis or, in adult individuals, from the observation of secondary sexual characters (e.g. length of the tail and of the claw in the front flipper). Otherwise, specify by NI (for Not Identified).

#### **13.4.** Protocol for dead sea turtles

#### Necroscopy

The carcass should be placed on its back, trying to wedge it with an object so that it does not wobble from side to side. The plastron should be removed and separated from the carapace through an incision on the outside edge (yellow line). The incision should be made with special attention, with the use of a short blade or by cutting with a horizontal tilt to avoid affecting the integrity of the interior organs.

Once the inside of the plastron is accessed, cut the ligament attachment to the pectoral and pelvic girdle to pull back the plastron and reach the muscles and then the internal organs.

Qualitative evaluation of the trophic status of the animal should be made, including the atrophy of pectoral muscles (none, moderate, severe), fat thickness in joint cavities and on coelom membrane (abundant, normal, low or none).



**Figure 13-4.** Sequence of turtle necropsy: a) Ventral view of a dead turtle. Yellow line indicates the way to separate the plastron from the rest of the turtle; b) Horizontal cuts to prevent affecting the interior organs; c) Ventral view of the opened turtle (fat reserves (brown) can be observed on the muscles). (From UNEP/MAP 2019).

# Tissue collection

Before sampling the content of the gastrointestinal tract, collect 10 g each of the following tissues for contaminants analysis, which should be wrapped in aluminium paper and kept at -20 °C:

- Muscle
- Liver
- Subcutaneous fat from different body parts
- Kidney
- Carapace scutes

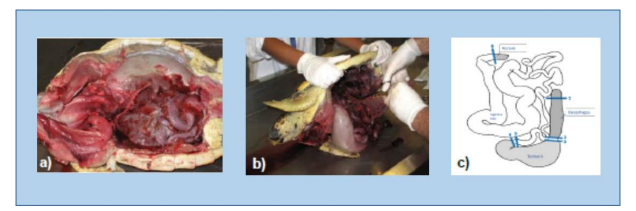
Each tissue stored in aluminium foil must be labelled with the standard identification code of the animal.

In case of turtles dead in rescue centres (max 1-2 h after death), collect:

- Epidermis for analysis of biomarkers and contaminants: take 10-20 g from the neck and forelegs preserved in aluminium foil, store in liquid nitrogen or dry ice.
- Liver for biomarker and contaminant analyses: 10-20 g wrap in aluminium paper and store in liquid nitrogen or dry ice.
- Blood for contaminants: 5-10 ml in tubes and store at -20 °C.

#### Gut content analysis

**Extraction of the gastrointestinal system**: expose the gastrointestinal system (GI) by removing the pectoral muscles and the heart of the animal. The blood can be emptied from the abdominal cavity by carefully rolling the turtle onto a side. Clamp the oesophagus proximal to the mouth and clamp the cloaca, the closest to the anal orifice. Remove the entire GI and place it on the examination surface. Isolate the different portions of GI (oesophagus, stomach, intestines) by strangling and cutting between the 2 clamps (see the blue solid lines) the gastro-oesophageal sphincter and the pyloric sphincter.



**Figure 13-5.** Sequence of extraction and preparation of sections of the digestive tract (GI) a) Remove the pectoral muscle and the heart; b) Extraction of the GI; c) Sketch of the entire GI. Blue lines indicate where clamps must be attached in order to separate the 3 different GI sections (Drawing by V. Hergueta) (From UNEP/MAP 2019).

**Note the external lesions** of the GI that can be attributed to litter. Before opening up the digestive tube, examine the outer wall to observe possible perforations by foreign bodies or areas of necrosis. Also note secondary lesions, particularly a peritonitis following on a perforation of the digestive tube, an invagination of the digestive tube, an occlusion, etc. Photograph every lesion observed, taking care to get an overall view as well as close-up (macro-lens) photographs.

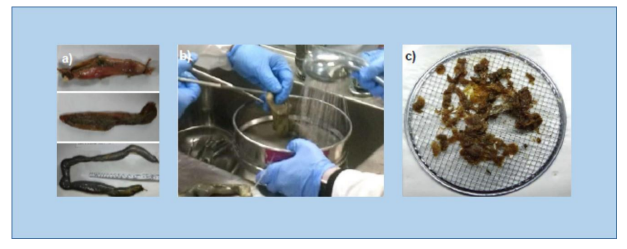
The three parts of the gastrointestinal system (i.e. oesophagus, stomach, intestines) should be removed by adding a second strangling at the cut edge to prevent spillage of the content.

Each GI section should be opened lengthways using a scissor and slide the material directly out of the section onto a 1 mm mesh sieve. The content should be cleaned with current and abundant tap water to remove the liquid portion, the mucus and the digested unidentifiable matter.

The content should be inspected for the presence of any tar, oil, or particularly fragile material, which should be subsequently removed and treated separately.

All material should be rinsed, collected in the 1 mm sieve, and placed in tubes or in zipped bags, reporting the sample code (individual code, respective GI section) and stored at -20 °C, pending the laboratory analyses.

For the separation of macrolitter and microlitter, the material should be slid out of the section directly onto a 5 mm mesh sieve superimposed on a 1 mm mesh sieve. Then, proceed with the rinsing and the storing of the material collected as described above, for both 1 and 5 mm sieves, reporting the samples code (individual code, respective GI section and size class (>5mm or 1-5mm)).



**Figure 13-6.** Digestive tract analysis: a) Separated GI sections: Oesofagus (up), stomach (middle) and intestines (down); b) Section opening and gut content lavage; c) Gut content extracted. (From UNEP/MAP 2019).

# 13.5. Protocol for live sea turtles

Live sea turtles hospitalized in rescue centres should be manually removed from water for the 30 min sampling period. The cares and procedures carried out on the rescued turtles for all the rehabilitation period should be performed in accordance with routine veterinary practices and guidelines for the conservation and rehabilitation of marine turtles. All the biological samples collected will be used for biomarker and chemical analyses.

#### Tissue collection

The collection of biological tissues such as blood and skin biopsy must be made with the support of the centres' veterinary while faeces can be collected by the volunteers or the operators of the rescue centre. Each tissue, stored in aluminium foil or Eppendorf, must be labelled with a unique ID for each individual.

#### **Blood sampling**

Blood samples (2-6 mL) are to be obtained from the dorsal cervical sinus using a disposable syringe. A small amount should be used for blood smears and the rest transferred into solvent-rinsed glass vials (10 or 5 ml) with Teflon caps containing heparinized saline (heparin sodium) following gently mixing of the tubes.

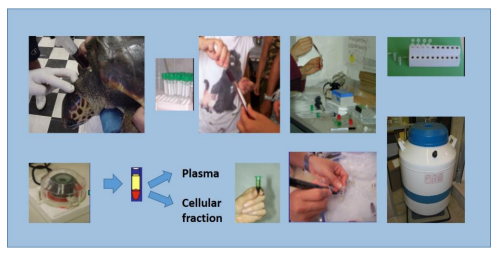


Figure 13-7. Blood collection and procedure for its processing and conservation.

A small amount of blood (two drops) is used for each animal to obtain two blood smears. Two blood smears for each sample are prepared in double (two slides). Once the blood is taken using a syringe, a drop of blood must be transferred to each slide. The smear of blood is done using a third clean slide as shown in the picture. Allow the blood film to air-dry. Slide fixing should be done the same day of sampling, after the slide is completely dry. Immerse the slides in ethanol for 10 minutes. Allow the blood film to air-dry. Place the slides in the appropriate slide boxes for further analysis.

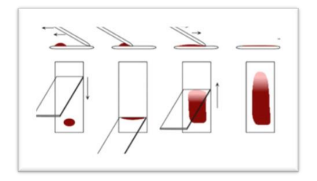


Figure 13-8. Blood smear technique explained in 4 steps.

A part of the blood (2 - 5 ml) is transferred into smaller (1.5 ml) centrifuge tubes and centrifuged at 5000 x g for 5 minutes for the separation of plasma that is immediately transferred into smaller plastic tubes (0.5 ml) containing a small amount of antiprotease cocktail (5  $\mu$ l) and stored into dry ice or liquid nitrogen (make a small hole in the upper part of the tubes to avoid the break when taking them out of the liquid nitrogen).

A part of the whole blood (1 ml) is stored without centrifugation in plastic tubes in liquid nitrogen or dry ice or -20°C.

500 uL of whole blood and 500 uL of mixture (RPMI and DMSO conservation mixture, 80:20) will be transferred into smaller (1.5 ml) centrifuge tubes and placed into liquid nitrogen or dry ice for biomarker analysis (comet assay).

# Skin biopsy sampling

Skin biopsy is performed (eventually after local anaesthesia and disinfection of the skin) using sterile iron punches. The dimension of the punches (diameter: 4 or 6 or 8 mm) depends on the weight of the animal. Once collected, the biopsy must be stored in aluminium foil and stored immediately in liquid nitrogen for enzymatic, cellular and molecular biomarkers. If the liquid nitrogen is not available, the biopsy can be stored in RNA-Later in an Eppendorf at room temperature for 24 hours and after at 4 °C.

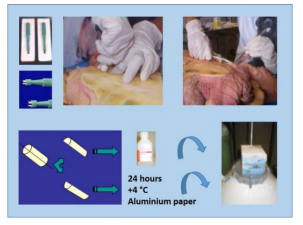


Figure 13-9. Skin biopsy removal and storage of the samples.

# Carapace sampling

A small amount of the superficial part of the carapace (free of epibionts, barnacles or algae) is to be removed by using a sterile scalpel; from several scutes, approximately 0.25 g of tissue (about 1-2 mm thick) are carefully removed using a disposable scalpel with a plastic handle and a stainless steel blade, by moving parallel to the carapace surface. Only the most superficial keratinous layer must be taken without penetrating the keratinous layer-bone interface below. Scute scrapings taken from carapace will be stored in aluminium paper at - 20  $^{\circ}$ C.



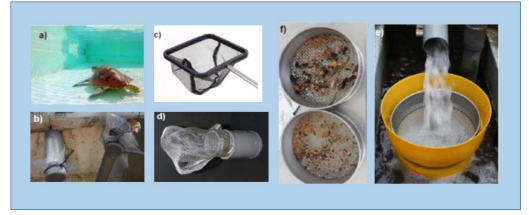
Figure 13-10. Carapace collection procedure and storage of the samples.

# Faeces collection

From the day that the animal arrives at the rescue centre, the excreta must be collected every day (or whenever excreta are expelled) for 2 months. The collected faeces will be analysed only for the individuals remaining at least 1-month minimum in the rescue centre. The collection method is as follows:

- Control the water tank daily by filtering through the 1 mm mesh sieve according to the following methods: collect the faeces manually with a 1 mm mesh dip net or put a 1 mm mesh flexible collector in the drain tube or place a 1 mm mesh rigid sieve under the drain
- Collect and store excreta and plastic in the same tube
- Use plastic containers (phthalate free) for collection
- Seal the test tube with the code, store the tube at -20 °C

Collect the excreta for the entire period of hospitalization.



**Figure 13-11.** Sequence of faeces sampling. A) The turtle is disposed in an individual tank; b) A 1mm mesh sieve is disposed in discharge tubes; c) A 1 mm dip net for handling faeces; d) Collector with 1 mm mesh disposed in discharge tube for filtering water tank; e) An 1 mm mesh rigid sieve down discharge tube for filtering water tank; f) Sample collected in a rigid sieve (from MAP/UNEP 2019).

# 13.6. Litter analysis and classification

#### Macrolitter detection

- Sort prey and/or litter items into separate categories under a stereomicroscope, taking care of recording their weight.
- Measure the size of litter items and classify litter categories.

In addition, the following parameters should be recorded:

- For all categories: the dry mass (grams, precision 0.01 g) of each category; dry the sample at room temperature for 24h minimum or in a stove at 35 °C for 12 h.
- For litter categories only: the number of fragments and items in each category; a fragment is a piece of litter that can be identified while an item is a set of fragments that seem to originate from the same piece of litter.
- For the plastic litter categories only: the total number of plastic fragments per colour category, with specifics as follow:
  - Total number of white-transparent plastic fragments.
  - Total number of dark coloured plastic fragments (black, blue, dark green...).
  - Total number of light-coloured plastic fragments (cream, yellow, pink, light green...).
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

#### Microlitter detection

- Examine the fraction 1-5 mm in the Petri dish under a stereomicroscope for particles resembling microplastics. Cover the petri dish with glass lids during observation not to contaminate the sample.
- Photograph, count and record the type, colour and maximum length of microplastic particles using image analysis software.
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

The limit detection for MSFD is 1 mm. Building on the findings of the testing phase of the Plastic Busters MPAs project, it's recommended to also examine the fraction 0.1-1 mm in stranded organisms.

# Litter categories

Categorize marine litter according to the categories showed in Table 13-1. The categorization of gastrointestinal tract contents and excreta is based on the general "morphs" of plastics (sheet-like, thread-like, foamed, fragment, other) or other general rubbish or litter characteristics. This is because in most cases, particles can't be unambiguously linked to particular objects. But where is possible, under notes in datasheets, the items should be described and assigned a litter category number using the "Joint List" developed by the TSG ML group (Fleet et al., 2021). In addition, it is important to measure and quantify also natural items (food and/or no food).

**Table 13-1.** Classification of Marine Litter items plus Food remain and Natural no food remain (fromINDICIT 2018).

ТҮРЕ	CODE	DESCRIPTION
Industrial Plastic	IND PLA	Industrial plastic granules usually cylindrical but also sometimes oval, spherical or cubical shapes.
Use sheet	USE SHE	Remains of sheet, e.g. from bag, cling-foil, agricultural sheets, rubbish bags
Use thread	USE THR	Threadlike materials, e.g. pieces of nylon wire, net-fragments, woven clothing
Use foam	USE FOA	All foamed plastics e.g. polystyrene foam, foamed soft rubber (as in mattress filling)
Use fragment	USE FRAG	Fragments, broken pieces of thicker type plastics, can be a bit flexible, but not like sheet like materials
Other use plastics	USE POTH	Any other plastic type of plastics, including elastics, dense rubber, cigarette filters, balloon pieces, soft airgun bullets Specify in the column "Notes".
Litter other than plastic	OTHER	All non-plastic rubbish and pollutant
Natural food	FOO	Natural food for sea turtles (e.g., pieces of crabs, jellyfish, algae)
Natural no food	NFO	Anything natural, but which cannot be considered as normal nutritious food for sea turtle (stone, wood, pumice, etc.)

# Collection of data

For each organism, an assessment is made of:

- 1. Frequency of occurrence (%) of ingested macro and microlitter for each species, calculated as the percentage of the individuals examined with ingested macro- and microplastics.
- 2. Abundance (N) of macro and microlitter ingested per individual (average number of items/individual) for each species, calculated as a total and per category. Since currently there are inconsistencies in the literature in reporting abundance of ingested litter, it is recommended to report average number of items per individual considering both all individuals examined and only individuals found with ingested macro and litter.
- 3. Total dry weight (g) of the detected waste expressed on grams (precision: second decimal place). This weight refers to each single category found in a specific organ (or faeces) of the specimen.

Other information as colour of items, polymer of the different items (at least 10% of the total items) and different incidence of litter in oesophagus, stomach and intestine, incidence and abundance are useful for research and impact analysis.

#### 13.7. Analysis of plastic tracers and PBTs

#### Plastic additives

The compounds to be detected in different tissues/fluids are:

- <u>*Phthalates:*</u> a group of chemicals widely used as additives to make plastics more flexible and harder to break; they can interfere with endocrine system (Baini et al., 2017).
- <u>Bisphenol A</u>: used in the production of polycarbonate, can have endocrine disrupting effects (Crain et al., 2007; Halden, 2010; Oehlmann et al., 2009) and the styrene and polyvinyl chloride monomer, used in the production of polystyrene and polyvinyl chloride (PVC), can be carcinogenic and/or mutagenic (Lithner et al., 2011; Papaleo et al., 2011; Xu et al., 2004).
- <u>Polybrominated diphenyl ethers</u>: they belong to the group of brominated flame retardants (BFRs), which are used in various polymeric materials such as plastic parts, resins, textiles, and other substrates to reduce their fire hazards (BSEF 2003; Król et al. 2012).

#### Persistent, bioaccumulative and toxic substances (PBTs)

In addition to the plastic additives that may leach from plastics when released into the marine environment, plastics also tend to adsorb in their surface persistent bioaccumulative and toxic substances (PBTs) (e.g. organochlorine compounds OCs, PAHs and PBDEs) and metals (e.g., lead, copper and cadmium) that are present in the seawater.

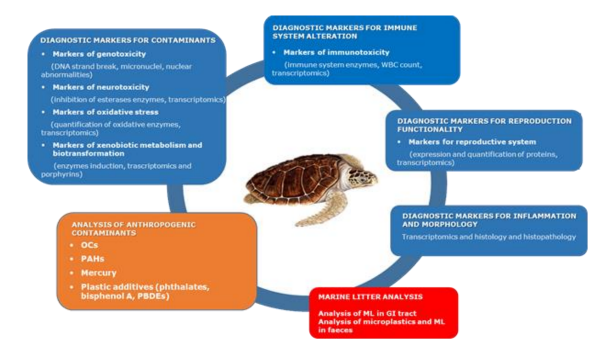
Depending on the compounds and the tissue to be analyzed, different methods should be applied to detect the presence of plastic-related contaminants in the sea turtles (Annex IV).

#### 13.8. Biomarkers analysis

The toxicological effects associated with the presence of marine litter can be evaluated using a set of diagnostic and prognostic methodologies, by means of biomarkers. A non-exhaustive list of existing biomarker approaches and plastic tracers' contaminants that are usually applied in sea turtle analyses is reported in Annex IV.

Biomarkers have been selected on the basis of the level of biological responses and in relation to the main effects related to marine litter/microplastics ingestion. The selected biomarkers can diagnose different impacts related to: a) physical damages/effects of marine litter, b) exposure to/effect of chemical tracers, and c) exposure to/effect of adsorbed chemicals.

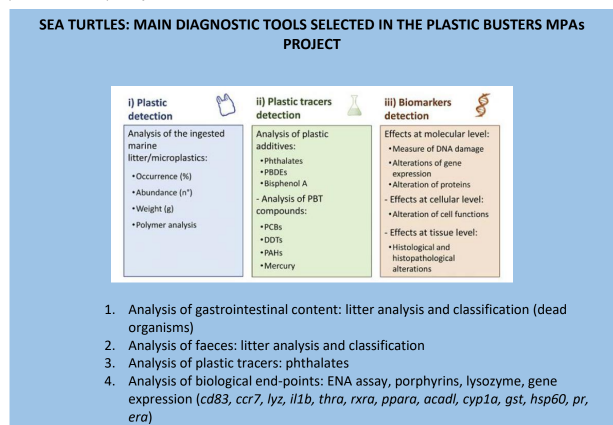
#### Set of protocols on harmonized marine litter monitoring approaches



*Figure 13-12.* A three-fold approach to detect the marine litter presence and impacts to sea turtle species (dead and live).

Starting from this initial list and building on the findings of the testing phase of the Plastic Busters MPAs project, the most suitable diagnostic tools to detect the presence and impact of ML on sea turtles are proposed here below.

**Table 13-2.** Main diagnostic tools selected in the PLASTIC BUSTERS MPAs project to detect the presence and impact of ML in sea turtles.



#### 13.9. Materials & Equipment for sampling

- Boots
- Camera
- Clamps (at least 6) or roast wire
- Clips with claws
- Containers for samples (Bottle/zipped bags)
- Cooler
- Cut-resistant gloves
- Garbage bag
- Glasses and protective mask or shield
- Nitrile Gloves
- Integral protective suit
- Measuring decimetre
- Measuring tape
- Metal containers
- Metal spoon
- Pen
- Permanent marker
- Sampling sheets
- Scalpel
- Scissors
- Sieve with 1 mm mesh

- Sieve with 5 mm mesh
- Transport bins or containers
- Aluminium foils
- Cryoboxes
- Cryovials
- Eppendorf (0.5 ml. 1.5 ml. 2.0 ml)
- Falcon tubes
- Liquid nitrogen Dewar (in alternate dry ice)
- Paper and block-notes
- Paper towels
- Pasteurs
- Pencils
- Plastic Sealable bags
- RNA-Later
- Ruler
- Scalpels
- Thermic bags
- Tweezers

# References

UNEP/MAP, 2019. Protocols for Monitoring Interactions between Marine Litter and Marine Turtles (Ingestion and entanglement) with a View to Harmonize Methods of Data Collection for Monitoring and Assessment in the Mediterranean. Document UNEP/MED WG.464/6.

Guidance on Monitoring of Marine Litter in European Seas, 2022. A guidance document within the Common Implementation Strategy for the marine Strategy Framework Directive - Marine Litter Impact on Biota. MSFD Technical Group on Marine Litter - draft.

INDICIT II, Final report, 2021. Deliverable D1.6, of the European project "Implementation of the indicator of marine litter on sea turtles and biota in Regional Sea conventions and Marine Strategy Framework Directive areas".

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

# 13.10. Sampling & recording sheets

# Monitoring Marine Litter in stranded sea turtles

Species:	ID code:				
Location/Country:	Latitude		Longitude		
Location, country.					
Discovery circumstances	□ By catch/Fishery	🗆 At sea	🗆 Stran	ded	
Discovery circumstances	🗆 Dead RC 🛛 🗆 Ot	her 🗆 Unknowi	n		
Cause of mortality (Please specify according to the					
toolkit) Date of discovery					
Date of discovery					
Date of necroscopy					

Animal body condition							
Conservation status	🗆 Level 1	🗆 Level 2	🗆 Level 3		Level 4		
Healt status (plastron shape)	🗆 Poor (concave	2)	Fair (plane)	🗆 Good	(convex)		
Main injuries	<ul> <li>No injuries</li> <li>Abrasion</li> </ul>	🗆 Fractu	re 🗌 Amput	ation	□ Sectioning		
Affected parts	□ Wings Other	□Tail	□Neck	$\Box$ Head			
Fat reserve	🗆 Thin	🗆 Fat	🗆 Normal	🗆 Not r	ecorded (NR)		

	Collected tissues	N°. of aliquots
Muscle		
GI tract		
Liver		
Fat tissue		

Picture 🗌

**Picture ID:** 

Necropsy performed by:

Name and Institution:

Biometric measurement				
1. CCLst	cm	6. HW	cm	
2. CCWst	cm	7. PTL	cm	
3. SCLst	cm	8. CTL	cm	
4. SCWst	cm	9. CaCL	cm	
5. CPL	cm			
Weight (kg):				
Sex:  Male Identified	🗆 🗆 Fema	ale 🗆 No	ot	

# Note and remarks:

# Monitoring Marine Litter in live sea turtles

Species:			ID code:		
Date and Time: Arrival			Sampling		
Location/Country:					
Rescue site:		Latitude	Longitude	Rescue centre:	
Health status		1			
Cause of mor	<b>bidity</b> ording to the toolkit)				
Picture ref.:					
Carapace Len	gth	CCL (cm):		CCW (cm):	
Aprox. Age		Adult Sub-adult Juvenile			luvenile
Weight (kg):		Sex:  Male Female Not Identified			Over Identified
Marine Litter		<ul> <li>Entanglement</li> <li>Presence in GI tract</li> <li>None</li> </ul>			
		Liquid N <sub>2</sub>	RNA later	DMSO	Cell medium
	Whole blood				
	Blood smears				
	Plasma				
Aliquots	DMSO:RPMI conservation mix				
	Excreta				
	Carapace				
	Adipose tissue				
	Skin biopsy				
Slices	Treatment	Dose	Time	Hour-notes	

Excreta collection				
Date	N°. aliquot stored at -20 °C			

Note and remarks:

# **ANNEX IV**

	CHEMICAL COMPOUND	TISSUE/SAMPLE	ANALYTICAL METHOD
PLASTIC ADDITIVES	Phthalates	Blubber, muscle, liver	Baini et al., (2017), Fossi et al., (2016), Savoca et al., (2018)
		Blood	Takatori et al., (2004), Notardonato et al., (2021)
	Bisphenol A	Muscle	Ballesteros-Gómez et al., (2009),
		Blubber, liver	Xue et al., (2016), Guerranti et al., (2014), Di Renzo et al., (2021)
		Blood	Cobellis et al., (2009)
	Polybrominated diphenyl ethers	Blubber, muscle, liver, blood	Muñoz-Arnanz et al., (2016), Guerranti et al., (2014)
ADSORBED CONTAMINANTS	Polycyclic aromatic hydrocarbons	Blubber, muscle, liver, blood	Marsili et al., (2001), Cocci et al., (2018)
	Organochlorine contaminants	Blubber, muscle, liver, kidney, brain, blood	Marsili and Focardi, (1997), Cocci et al., (2018), Gómez-Ramírez et al., (2020)
	Mercury	Blood, skin, muscle, kidney, liver	Correa et al., (2013), Gómez-Ramírez et al., (2020)

 Table A-5. Tissues and methods to be used to detect plastic tracers in sea turtles.

**Table A-6.** Effects measured in sea turtles by the biomarker approach. The analysis on brain, liver, kidney and muscle can be performed only in dead sea turtle (level 1 - fresh carcass).

EFFECT	TISSUE	TEST
GENOTOXICITY	Blood	Comet assay (Molino et al., 2019) (*) Mn test (Bolognesi et al., 2006) ENA assay (Bianchi et al., 2022); (Pacheco and Santos, 1997)
OXIDATIVE STRESS	Plasma, skin	LPO (Fossi et al., 2016), Casini et al., 2018) CAT (Fossi et al., 2013) Cat, gpx, sod (Coccie et al., 2019)
IMMUNOTOXICITY	Blood	Total and differential white blood cells (WBC) count (Casal and Orós, 2007; Davis et al., 2008; Caliani et al., 2019) H:L ratio (Caliani et al., 2019) Respiratory burst (Secombes, 1990; Caliani et al., 2019; Bianchi et al., 2022) TAS assay (Miller et al., 1993; Caliani et al., 2019; Bianchi et al., 2022) Lisozyme enzyme (Keller et al., 2006; Caliani et al., 2019; Bianchi et al., 2022) casp8, casp9, TRAF (Karami et al. 2017; Mathieu-Denoncourt et al., 2015) (*)
	Plasma	CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015) (*) Vitellogenin (Fossi et al., 2004)
REPRODUCTION	Plasma, skin	Vitellogenin (Herbst et al., 2003) CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*) ERa (Cocci et al., 2018)
HISTOPATHOLOGY INFLAMMATION AND MORPHOLOGY	Liver	Histopathology, histology (Pedà et al. 2016; Karami et al. 2017; Batel et al., 2018) (*)
XENOBIOTIC METABOLISM AND BIOTRANSFORMATION	Blood, skin, excreta, liver	CYP1A; AHR, CYP3A (Fossi et al. 2014, Panti et al. 2011; Rochman et al., 2013) (*) Porphyrins (Guerranti et al., 2014) (*) Cyp1a, Cyp1b, gstt1 (Cocci et al 2018, 2019)
NEUROTOXICITY	Brain, muscle, plasma	AChE, BChE (Casini et al., 2018) (*)
CELLULAR STRESS	Blood, skin, liver, kidney	PPARA, PPARG, HSP70, GPX, E2F1 (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*) Gamma glutamyl transferase (GGT) (Nematdoost Haghi and Banaee, 2017) (*)

Cortisol and corticosterone (Flower et al., 2015) LDH (Nematdoost Haghi andBanaee, 2017) (*) HSP70, HSP90 (Cocci et al 2018)

(\*) effects detected after laboratory or field exposure with MPs or plastic-related contaminants.



# 14 Methodology for monitoring presence and effects of marine litter in seabirds

This document describes the methodological approach for monitoring the presence and effects of Marine Litter in seabirds, which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML, and on the results of the project's testing phase.

# PREPARED BY

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



#### 14.1. Species sampling

The presence and impact of marine litter in Mediterranean seabirds (*Calonectris diomedea, Puffinus yelkouan, Ichthyaetus audouinii*) can be investigated and/or monitored in animals dead, on beaches or from accidental mortalities such as long-line victims. Live animals could be sampled in the rescue centres during the hospitalization or in their colonies. In case of doubt about the species identification, refer to identification guide (e.g. www.cites.org) or an expert in the field.

The seabirds above indicated are protected species, therefore only authorized people can handle live and dead animals or parts of them. Upon finding the animal, its management and recovery should be reported and coordinated with the responsible Authorities. Note that a CITES permit is asked if a specimen or sample has to be sent/received.

#### 14.2. Data to be recorded

Taking pictures of the animal before handling it is key to verify the circumstances of the finding and to *a posteriori* confirm or clarify the noted information if doubts or difficulties are encountered in identifying the species, the lesions, the state of the individuals and the elements responsible for the interaction.

All the sea birds' data should be noted down in a sampling sheet (see paragraph 14.11).

# 14.3. Conservation/health status of the organism

With regards to the health status of the organism, two cases are possible: the seabird may be live, or dead. Overall, 5 different situations can be observed:

- Level 1: the animal is live. In this case biological samples (blood, plasma, faeces) can be removed for biomarkers and chemical analyses. In animals that have just died (< 2 hours post mortem), GI is adequate for litter ingestion analysis and other tissues (muscle, liver) can be used for biomarker and chemical analyses.
- Level 2: Fresh carcass (< 24 hours post mortem), adequate for litter ingestion analysis from necropsies and chemical analysis.
- Level 3: Moderate decomposition, adequate for litter ingestion analysis from necropsies and chemical analysis.
- Level 4: Advanced decomposition, adequate for litter ingestion analysis from necropsies and chemical analysis.
- Level 5: Mummified or skeletal remains. In this level individuals have usually lost their gastrointestinal material and thus, the analysis of litter ingestion is not possible.

#### Discovery circumstances

Note the circumstances among the 2 categories:

- Stranding: animal found stranded on the beach or in the shoreline.
- Dead at the recovery centre: the animal arrived live but died during its hospitalization.

#### Possible cause of morbidity and mortality, type of impact

If possible, the type of interaction with human activities and impact observed or suspected on dead or live stranded individuals should be deduced from external or organs observations during the necropsy and complemented with veterinarian examinations. Also, an inspection of the oral cavity should be conducted for the presence of foreign material. Then a choice among the following categories should be made and the notes and remarks box should be completed with the help of the pathologist (if this is requested):

- Bycatch/Fisheries related: ingested hook, individual trapped in a fishing gear, individual drowned in a fishing gear.
- Entanglement in litter: entanglement in litter other than related to fishing activity. Please fill the column "Entanglement type" and "Litter causing entanglement".
- Ingestion of litter: digestive obstruction or occlusion, perforation, or other impacts.
- Anthropogenic trauma.
- Natural trauma or natural disease.
- Oils: Ingestion or external impregnation with oils.
- Unidentified: Impossible to know the cause of death/stranding, no remarkable damages, injury or disease.
- Other: Please specify in the column "Notes".

#### Main injuries

In case of injuries, the main type of injury (fracture, amputation, sectioning, abrasion or other) and the affected body part should be reported.

If the individual has been found entangled in litter, the type of material in which the seabird was found should be specified.

#### Biometric Measurements

Several basic and body lengths (Culmen Lenght -CL, Wing Length -WL, Head Length -HL, Bill Depth -BD, Weight, Tarsus Length -TL) can be measured (in centimetres, precision 0.01 cm), as well as the weight (in kilograms, precision 0.01g). If possible, the sex (male or female) should be noted, which is determined by gonads observation. Otherwise, specify by NI (for Not Identified). Age, the only variable found to influence litter quantities in stomach contents, is largely determined based on development of sexual organs (size and shape) and presence of *Bursa of Fabricius*.

# 14.4. Protocol for dead seabirds

#### Dissection procedure

Immediately after sampling, label the animal (unique ID for each individual) and transport it to the laboratory in ice containers and store at -20 °C until dissection for litter and contaminant analysis. Thaw the animal in the laboratory at room temperature, then dissect out gastrointestinal tract (GI) for litter analysis and the other tissues for contaminant analyses.

# Tissue collection

For contaminants assessment, about 10g of each of the following tissues should be collected, wrapped in aluminum foil and stored at -20 °C: muscle, liver, subcutaneous fat from different parts of the body, kidney. Each tissue stored in aluminium foil must be labelled with the standard identification code of the animal (unique ID for each individual).

#### Gut content analysis

The stomachs of dissected birds are to be opened by scissors or scalpel. Stomach contents should be carefully rinsed in a sieve with a 1 mm mesh and then transferred to a petri dish for sorting under a

binocular microscope. The 1 mm mesh is to be used because smaller meshes become easily clogged with mucus from the stomach wall and with food-remains.

If oil or chemical types of pollutants are present, these may be sub-sampled and weighed before rinsing the remainder of the stomach content. If sticky substances hamper further processing of the litter objects, hot water and detergents should be used to rinse the material clean as needed for further sorting and counting under a binocular microscope.

# 14.5. Protocol for live seabirds

The sampling of live seabirds can be applied in seabird colonies (free-ranging animals) or in animals hospitalized in rescue centres.

In seabird colonies, nests can be difficult to access. Safety requirements for boating, climbing and hiking should be followed. In some risky conditions, despite protocols being simple, only experts should be asked to take samples.

Moreover, seabird welfare and safety should be a priority for coordinators and operators, and unnecessary stress to birds should be avoided. Some precautions, such as cover bird head, avoid noise, exclude from sampling nests in unfavourable conditions, and fast sampling procedures should be considered case by case.

# Sample collection

- The collection of biological samples on live organisms should be made by authorized personnel.
- Biological tissues (blood, oil gland secretion, faeces, and abandoned eggs) must be collected, processed, and immediately stored in liquid nitrogen or dry ice. Each tissue must be stored in aluminum foil and labelled. All the biological samples collected are to be used for biomarker and chemical analyses.

# Blood sampling

- Blood (the amount depends on the size of the animal ranging from about 50 ul to 2 ml) should be collected from a brachial vein using an insulin syringe. The brachial/ulnar vein is located just beneath the ventral surface of the humeral-radialulnar joint. Extend the wing, possibly with the aid of a collaborator, and clear the area of feathers around the ulnar-humoral joint using a cotton ball soaked in distilled water until the brachial vein is visible.
- Then, with a 23-25-gauge needle (internal diameter of about 0.3 mm) kept in orthogonal sense with respect to the vein, gently prick. If the blood flows poorly, insert the needle slowly into the vein and use a needle bore size more appropriate.
- Once the blood is flowing, remove the needle and use the syringe for collecting it (Owen, 2011). To stop the blood from flowing, press on the puncture site using cotton wool for half a minute. Allow the wing to fold naturally against the body, securing in that position to prevent flapping. The blood should be transferred into a solvent-rinsed glass vials (10-5 ml) with Teflon caps containing heparinized saline (heparin sodium) and the tubes gently mixed.



**Figure 14-1.** Blood collection from the brachial vein in a specimen of Calonectris diomedea. (Photo ©UNISI).

One drop of blood is enough to obtain a blood smear. Each sample must be done in double. Once the blood is collected using a syringe, a drop of blood must be transferred to each slide. The blood smear should be performed by a different operator from who makes the blood collection, as blood immediately coagulates and, contemporary, bleeding must be stopped. The blood smear is done using a third clean slide as shown in the picture. Dry the slides at the air. Slide fixing shall be done the same day of sampling, after the slide is completely dry. Immerse the slides in ethanol for 10 minutes and dry the slides in the air, then place the slides in the appropriate slide box.

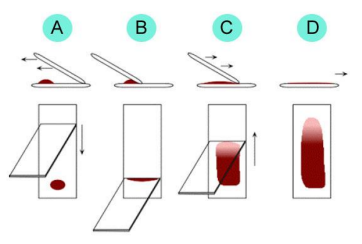


Figure 14-2. Blood smear collection.

A part of the blood (2 ml) is transferred into smaller (2 ml) centrifuge tubes and centrifuged at 5000 x g for 5 minutes for the separation of plasma and immediately transferred into smaller plastic tubes (0.5 ml) containing a small amount of antiprotease cocktail (5  $\mu$ l). Placed into ice dry or liquid nitrogen (make a small hole in the upper part of the tubes to avoid "explosion" when taking them out of the liquid nitrogen).

A part of the whole blood (1 ml) will be stored without centrifugation in plastic tubes in liquid nitrogen or dry ice or -20  $^{\circ}$ C.

500 uL of whole blood and 500 uL of mixture (RPMI and DMSO conservation mix, 80:20) will be transferred into smaller (2 ml) centrifuge tubes and placed into liquid nitrogen or dry ice for biomarker analysis (comet assay).

# Oil gland secretion sampling

This minimally intrusive protocol aims to collect a small quantity of oil from the uropygial gland of live birds, in order to detect contaminants (phthalates).

- Once the bird is kept in hand, gently massage the preen gland at the upper base of the tail. With bare hands, give a gentle squeeze after massaging the gland so that a small amount of oil can be obtained.
- The gland secretes a waxy substance, rather than a fluid oil as one could expect. Using a pair of metal tweezers that have not been in contact with plastic, remove a clean cotton wool from a glass jar.
- Gently massage the oil gland and wipe cotton wool over the gland 1-2 times to transfer the oil gland exudate to the cotton wool. Do this without touching latex gloves or other plastic items. Then, place the cotton wool back in a glass jar. Seal and label the jar (Hardesty et al., 2015).



Figure 14-3. The uropygial gland in a seabird (from CSIRO, 2013).

• After sampling, be careful with the glassware containing samples. To transfer samples from field to the laboratory, it may be useful to protect vials with packaging material, avoiding any plastic products even if vials are sealed. Use leather, corn or paper materials.

# Faeces sampling

Faeces can be taken into or next to each nest in the colony. Fresh faeces ejected during bird handling can be collected too. Dry, almost dry or fresh faeces should be collected using a teaspoon or spatula (about 1 g needed). Put them in a piece of aluminum foil which will be closed as an envelope (or in an Eppendorf). Beware that samples faeces are not contaminated with the soil or other external materials and do not mix excrement samples from different nests, or sites. After sampling, freeze samples at -20 °C (within 24-48 hours) to detect possible presence and effect of litter ingestion.

# 14.6. Litter analysis and classification

#### Macrolitter detection

- Sort prey or litter items from the bird stomach into separate categories under a stereomicroscope, taking care of recording their weight.
- Measure the size of litter items and classify litter categories.

In addition, the following parameters should be recorded:

- For all categories: the dry mass (grams, precision 0.01 g) of each category; dry the sample at room temperature during 24 h minimum or in a stove at 35°C during 12 h.
- For litter categories only: the number of fragments and items in each category; a fragment is a piece of litter that can be identified while an item is a set of fragments that seem to originate from the same piece of litter
- For the plastic litter categories only: the total number of plastic fragments per colour category, with specifics as follow:
  - Total number of white-transparent plastic fragments;
  - Total number of dark coloured plastic fragments (black, blue, dark green...);
  - Total number of light-coloured plastic fragments (cream, yellow, pink, light green...).
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

#### Microlitter detection

- Examine the content >1mm in the Petri dish under a stereomicroscope for particles resembling microplastics. Cover the Petri dish with glass lids during observation not to contaminate the sample.
- Photograph, count and record the type, colour and maximum length of microplastic particles using image analysis software.
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

The limit detection for MSFD is 1 mm. Building on the findings of the testing phase of the Plastic Busters MPAs project, it's recommended to also examine the fraction 0.1-1 mm.

# Litter categories

Categorize marine litter according to the categories showed in Table 14-1. The categorization of stomach contents is based on the general "morphs" of plastics (sheet-like, thread-like, foamed, fragment, other) or other general rubbish or litter characteristics. This is because in most cases, particles cannot be unambiguously linked to particular objects. But where is possible, under notes in datasheets, the items should be described and assigned a litter category number using the "Joint List" developed by the TSG ML group (Fleet et al., 2021). In addition, it is important to measure and quantify also natural items (food and/or no food).

**Table 14-1.** Categories for classification of items for sea birds (MSFD Protocol for the monitoring of litter ingested by seabirds, 2022).

	BIOTA c	ategor	ies for contents of digestive tract
PLA	PLASTIC	acronym	all plastic or synthetic items: note number of particles and dry mass for each category
IND	pellets	ind	industrial plastic granules (usually cylindrical but also oval spherical or cubical shapes exist)
	probab ind?	pind	suspected industrial, used for tiny spheres (glassy, milky,) (= microbeads)
	sheet	she	remains of sheet, eg from bags, cling-foil, agricultural sheets, rubbish bags etc
USE	thread	thr	threadlike materials, eg pieces of nylon wire, net-fragments, woven clothing; includes 'balls' of compacted material
	foam	foam	all foamed plastics, polystyrene foam, foamed soft rubber (as in matrass filling), PUR used in construction etc
	fragments	frag	fragments, broken pieces of thicker type plastics, can be bit flexible, but not like sheetlike materials
	other	Poth	any other, incl elastics, dense rubber, cigarette-filters, balloon-pieces, softairgun bullets, objects etc. DESCRIBE!!
RUB	OTHER RUBBISH	acronym	any other nonsynthetic consumer wastes: note number of particles and (in principle) dry mass for each category
	paper	рар	newspaper, packaging, cardboard, includes multilayered material (eg Tetrapack pieces) and aluminium foil
	kitchenfood	kit	human food remains (galley wastes) like onion, beans, chickenbones, bacon, seeds of tomatoes, grapes, peppers, melon etc
RUB	other rubbish	rubvar	other various rubbish, like processed wood, pieces of metal, metal air-gun bullets; leadshot, paintchips. DESCRIBE
	FISHHOOK	hook	fishing hook remains (NOT FOR HOOKS ON WHICH LONGLINE VICTIMS WERE CAUGHT - THOSE UNDER NOTES)
POL	POLLUTANTS (INDUS/CHEM WASTE)	acronym	other non-synthetic industrial or shipping wastes (number of items and mass per category (wet for paraffin)
	slag/coal	slag	industrial oven slags (looks like non-natural pumice) or coal remains
POL	oil/tar	tar	lumps of oil or tar (also note as n=1 and g=0.0001g if other particles smeared with tar but cannot be sampled separately)
102	paraf/chem	chem	lumps or soft mush of unclear paraffin, wax like substances (NOT stomach oil!); if needed estimate mass by subsampling
	featherlump	confea	lump of feathers from excessive preening of fouled feathers (n=1 with drymass) (NOT for few normal own feathers)
FOO	NATURAL FOOD	foo	various categories, depends on the species studied, and aims of study
NFO	NATURAL NON FOOD	nfo	anything natural, but which cannot be considered as normal nutritious FOOD for the individual

#### Collection of data

For each organism, an assessment is made of:

- 1. Frequency of occurrence (%) of ingested macro and microlitter for each species, calculated as the percentage of the individuals examined with ingested macro- and microplastics.
- 2. Abundance (N) of macro and microlitter ingested per individual (average number of items/individual) for each species, calculated as a total and per category. Since currently there are inconsistencies in the literature in reporting abundance of ingested litter, it is recommended to report average number of items per individual considering both all individuals examined and only individuals found with ingested macro and litter.
- 3. Total dry weight (g) of the detected waste expressed on grams (precision: second decimal place). This weight refers to each single category found in a specific organ (or faeces) of the specimen.

Other information as colour of items, polymer of the different items (at least 10% of the total items) and different incidence of litter in oesophagus, stomach and intentine, incidence and abundance are useful for research and impact analysis.

#### 14.7. Analysis of plastic tracers and PBTs

#### Plastic additives

The compounds to be detected in different tissues/fluid are:

- <u>*Phthalates:*</u> a group of chemicals widely used as additives to make plastics more flexible and harder to break; they can interfere with endocrine system (Baini et al., 2017).
- <u>Bisphenol A</u>: used in the production of polycarbonate, can have endocrine disrupting effects (Crain et al., 2007; Halden, 2010; Oehlmann et al., 2009) and the styrene and polyvinyl chloride monomer, used in the production of polystyrene and polyvinyl chloride (PVC), can be carcinogenic and/or mutagenic (Lithner et al., 2011; Papaleo et al., 2011; Xu et al., 2004).
- <u>Polybrominated diphenyl ethers</u>: they belong to the group of brominated flame retardants (BFRs), which are used in various polymeric materials such as plastic parts, resins, textiles, and other substrates to reduce their fire hazards (BSEF 2003; Król et al. 2012).

#### Persistent, bioaccumulative and toxic substances (PBTs)

In addition to the plastic additives that may leach from plastics when released into the marine environment, plastics tend also to adsorb in their surface persistent bioaccumulative and toxic substances (PBTs) (e.g. organochlorine compounds OCs, PAHs and PBDEs) and metals (e.g., lead, copper and cadmium) that are present in the seawater.

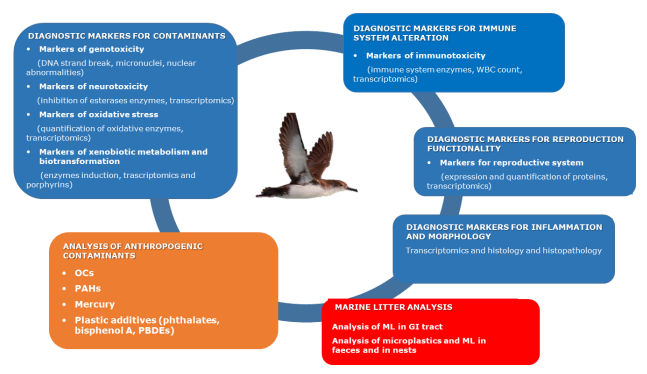
Depending on the compounds and the tissue to be analysed, different methods should be applied to detect the presence of plastic-related contaminants in the fish species (Annex V).

#### 14.8. Biomarkers analysis

The toxicological effects associated with the presence of marine litter can be evaluated using a set of diagnostic and prognostic methodologies, by means of biomarkers. A non-exhaustive list of existing biomarker approaches and plastic tracers' contaminants that are usually applied in seabirds analyses is reported in Annex V.

Biomarkers have been selected on the basis of the level of biological responses and in relation to the main effects related to marine litter/microplastics ingestion. The selected biomarkers can diagnose

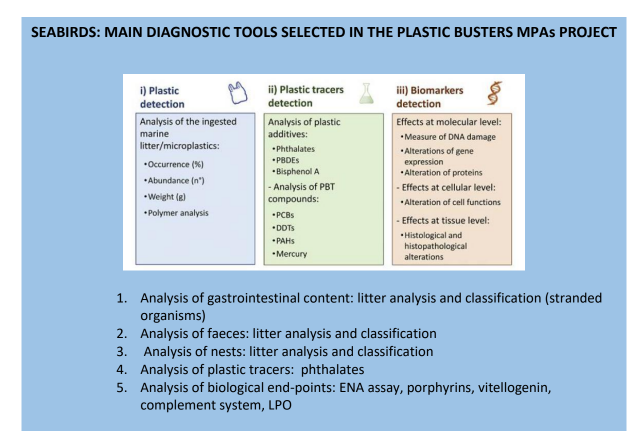
different impacts related to: a) physical damages/effects of marine litter, b) exposure to/effect of chemical tracers, and c) exposure to/effect of adsorbed chemicals.



**Figure 14-4.** A three-fold approach to detect the marine litter presence and impacts to seabirds species.

Starting from this initial list and building on the findings of the testing phase of the Plastic Busters MPAs project, the most suitable diagnostic tools to detect the presence and impact of ML on seabirds are proposed here below.

**Table 14-2.** Main diagnostic tools selected in the PLASTIC BUSTERS MPAs project to detect the presence and impact of marine litter in seabirds.



#### 14.9. Materials & Equipment for sampling

The following material and equipment are necessary for the correct application of the protocol:

- Boots
- Camera
- Clamps (at least 6) or roast wire
- Clips with claws
- Containers for samples (Bottle/zipped bags)
- Cooler
- Cut-resistant gloves
- Garbage bag
- Glasses and protective mask or shield
- Nitrile Gloves
- Integral protective suit
- Measuring decimetre
- Measuring tape
- Metal containers
- Metal spoon
- Pen
- Permanent marker
- Sampling sheets
- Scalpel

- Scissors
- Sieve with 1 mm mesh
- Sieve with 5 mm mesh
- Transport bins or containers
- Aluminium foil
- Cryoboxes
- Cryovials
- Eppendorf (0.5 ml. 1.5 ml. 2.0 ml)
- Falcon tubes
- GPS
- Liquid nitrogen Dewar (in alternate dry ice)
- Paper and block-notes
- Paper towels
- Pasteurs
- Pencils
- Plastic Sealable bags
- RNAlater
- Ruler
- Scalpels
- Spare batteries
- Thermic bags
- Tweezers

# References

Guidance on Monitoring of Marine Litter in European Seas, 2022. A guidance document within the Common Implementation Strategy for the marine Strategy Framework Directive - Marine Litter Impact on Biota. MSFD Technical Group on Marine Litter - draft.

Fossi, M.C, Vlachogianni, T., Anastasopoulou, A., Alomar, C., Baini, M., Caliani, I., Campani, T., Casini, S., Consoli, P., Cillari T., D'Alessandro, M., Deudero, S., Galgani, Galli M., F., Kaberi H., Panti, C., Pedà, C., E. Romeo, T., Scotti, G., Tsangaris, C., Zeri, C., 2019. Toolkit for monitoring marine litter and its impacts on biodiversity in Mediterranean MPAs. Interreg Med Plastic Busters MPAs project (D.3.3.2).

# 14.10. Sampling & recording sheets

# Monitoring Marine Litter in stranded/dead seabirds

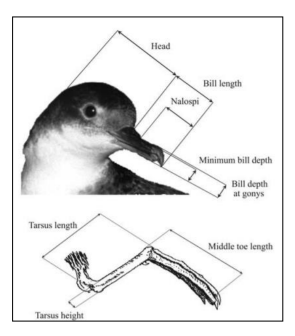
Species:		ID code:			
Ringing code:					
Location/Country:		Latitude	Longitude		
Discovery circumstances	□ Stranding □	Dead at recovery center			
Cause of mortality (Please specify according to the toolkit)					
Date of discovery					
Date of necroscopy					

Animal body condition								
Conservation status	🗆 Level 1	🗆 Level 2	🗆 Level 3	; [	□ Level 4			
Main injuries	<ul> <li>□ No injuries</li> <li>□ Abrasion</li> </ul>	□ Fracture	🗆 Amput	tation	□ Sectioning			
Affected parts	□ Wings	□Tail	□Neck	🗆 Head	🗆 Other			

	Collected tissues	N°. of aliquots
Muscle		
GI tract		
Liver		
Kidney		
Sub-cutaneous fat		

Necroscopy performed by:

Name and Institution:



Measurement							
Sex	🗆 Male	□ Male □ Female □ Not Identified					
<b>C</b> ulmen <b>L</b> ength (CL)			Bill Dept	th (BD)			
Wing Length (WL)			Weight	(kg)			
Head Length (HL)							

Note and remarks:

# Monitoring Marine Litter in live seabirds

Species:		ID code:			
Ringing code:		•			
Location/Country:		Latitude	Longitude		
Discovery circumstances		Dead at recovery cente	er		
Cause of morbidity (Please specify according to the toolkit)					
Date of discovery					

Measurement								
Sex	$\Box$ Male	🗆 Female	e 🛛 🗆 Not Ident	ified				
<b>C</b> ulmen <b>L</b> ength (CL)			Bill Depth (BD)					
<b>W</b> ing <b>L</b> ength (WL)			Weight (kg)					
Head Length (HL)								

	Collected tissues						
Whole blood Feathers							
Plasma		Uropygial gland					
Liver		Excreta					
DMSO:RPMI conservation mix		Egg					

# ANNEX V

	CHEMICAL COMPOUND	TISSUE/SAMPLE	ANALYTICAL METHOD
		Fat, muscle, liver	Baini et al., (2017), Fossi et al., (2016), Savoca et al., (2018)
	Phthalates Blood		Takatori et al., (2004)
		Oil gland secretion	Hardesty et al., (2015), Provencher et al., (2020)
PLASTIC ADDITIVES		Muscle	Ballesteros-Gómez et al., (2009)
	Bisphenol A	Fat	Xue et al., (2016)
		Blood	Cobellis et al., (2009)
	Polybrominated diphenyl ethers	Fat, muscle, liver, egg, blood	Muñoz-Arnanz et al., (2016), Sühring et al., (2022)
	Polycyclic aromatic hydrocarbons	Fat, muscle, liver, blood	Marsili et al., (2001)
ADSORBED CONTAMINANTS	Organochlorine contaminants	Fat, muscle, liver, blood	Marsili and Focardi, (1997), Sühring et al., (2022)
	Mercury	Blood, kidney	Correa et al., (2013), (Espín et al., 2012)

 Table A-7. Tissues and methods to be used to detect plastic-related contaminants in seabirds.

EFFECT TISSUE TEST Comet assay (Molino et al., 2019) (\*) GENOTOXICITY Blood Mn test (Bolognesi et al., 2006) ENA assay (Casini et al., 2018); (Pacheco and Santos, 1997) CAT, GST, LPO, GPX, GR, GSH (Yu et Liver, kidney al., 2018) (\*) **OXIDATIVE STRESS** LPO (Fossi et al., 2016), Casini et al., Plasma 2018) CAT (Fossi et al., 2013) Total and differential white blood cells (WBC) count (Casal and Orós, 2007; Davis et al., 2008; Caliani et al., 2019) H:L ratio (Caliani et al., 2019) Respiratory burst (Secombes, 1990; Caliani et al., 2019) ΙΜΜυΝΟΤΟΧΙCITY Blood TAS assay (Miller et al., 1993; Caliani et al., 2019) Lisozyme enzyme (Keller et al., 2006; Caliani et al., 2019) casp8, casp9, TRAF (Karami et al. 2017; Mathieu-Denoncourt et al., 2015) (\*) CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015) Plasma, Gonads (\*) Vitellogenin (Fossi et al., 2004) REPRODUCTION Vitellogenin (Herbst et al., 2003) CYP17A, CYP19, ERs, VTG, StAR Plasma (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (\*) Histopathology, histology (Pedà et HISTOPATHOLOGY INFLAMMATION AND Liver, kidney al. 2016; Karami et al. 2017; Batel MORPHOLOGY et al., 2018) (\*) CYP1A; AHR, CYP3A (Fossi et al. 2014, Panti et al. 2011; Rochman et **XENOBIOTIC METABOLISM AND** Blood, excreta, liver al., 2013) (\*) **BIOTRANSFORMATION** Porphyrins (Guerranti et al., 2014) (\*) AChE, BChE (Barboza et al., 2018) NEUROTOXICITY Brain, muscle, plasma (\*)

**Table A-8.** Effects measured in seabirds by the biomarker approach. The analysis on brain, liver, kidney and muscle can be performed only in dead sea birds (fresh carcass).

CELLULAR STRESS	Blood, liver, kidney	PPARA, PPARG, HSP70, GPX, E2F1 (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*) Gamma glutamyl transferase (GGT) (Nematdoost Haghi and Banaee, 2017) (*) Cortisol and corticosterone (Flower et al., 2015) LDH (Nematdoost Haghi and Banaee, 2017) (*)
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(\*) effects detected after laboratory or field exposure with MPs or plastic tracers.



This document describes the methodological approach for monitoring the presence and effects of marine litter in marine mammals, which has been developed within the framework of the Interreg Med Plastic Busters MPAs project, building on the most recent methodological advances of the MSFD TGML, Barcelona Convention CORMON, ACCOBAMS/ASCOBAMS, and on the results of the project's testing phase.

**PREPARED BY** 

# THE INTERREG MED PLASTIC BUSTERS MPAs PROJECT



# 15.1. Sampling approach

The ingestion of macrolitter and microlitter by Mediterranean marine mammals such as deep diver cetaceans species (*Physeter macrocephalus, Ziphius cavirostris*), coastal and pelagic odontocetes (*Tursiops truncatus, Stenella coeruleaolba, Delphinus delphis, Grampus griseus, Globicephala melas*), mysticete (*Balaenoptera physalus*) and pinniped species (*Monachus monachus*), and the potential related effects can be investigated and/or monitored in:

- **Dead organisms** which may have been stranded ashore, found at sea, etc.
- **Free ranging organisms** that have been sampled at sea.

Marine mammals are protected species, therefore only authorized people can handle live and dead animals or parts of them. Upon finding the animal, its management and recovery should be reported to and coordinated with the responsible authorities. Permits released by national competent authorities are required for the collection of cetacean biopsy samples. Note that a CITES (Convention on International Trade in Endangered Species of Wild Fauna and Flora) permit is required, if a specimen or sample has to be sent or received.

# 15.2. Protocol for dead animals

The protocol for the analysis of marine litter in stranded marine mammals was developed according to available protocols for other marine taxa (Lusher et al., 2017, 2018, Fossi et al 2018). The approach presented within this document, has already been integrated into a document enitled "*Best practice on cetacean post mortem investigation and tissue sampling*" (Ijsseldijk et al., 2019), developed jointly by ACCOBAMS and ASCOBANS.



Figure 15-1. A stranded cetacean. (Photo ©UNISI).

# Stage of decomposition

The stage of decomposition and the carcase quality is an important is an important determinant in subsequent analyses. Carcasses are assigned to one of five decomposition condition categories (DCC), determined by specific characteristics, as specified below:





Condition code 3



Condition code 2



**Condition code 5** 

Figure 15-2. Decomposition condition categories and associated codes.

- **CODE 1**: the animal is found live or dead at most by 2 h, adequate for *litter ingestion investigation, chemical analysis and biomarkers analyses.*
- Level 2 (Death within 24 h); normal appearance with minimal damage from scavenger animals; normal smell; minimal skin dehydration and rippling of the skin, and apparent mucous membranes; clean and shiny eyes; uninflated carcass, tongue and penis not protruding. Adequate for litter ingestion and chemical analyses.
- Level 3: Whole carcass, with evident swelling (tongue and penis protruding); skin not integrated with detachment areas; possible damage from scavenger animals; slight characteristic smell; apparent dry mucous membranes; eyes introflexed or missing. Adequate for litter ingestion and chemical analyses.
- Level 4: The carcass may be intact, but collapsed; wide areas of skin disepithelialization; severe damage from opportunistic animals; strong smell; muscles and blubbers easily removable and detachable from the bone; liquefaction of internal organs; allows to measure biometric data and assess the *presence/absence of ingested plastic and chemical analyses*.
- Level 5: Often with dehydrated skin and dry over the bones; completely dry; the analyses of litter ingestion or chemicals are not possible.

Table 15.1 provides guidance on sample collection and applicable analytical procedures, including contaminant analysis and ingestion of marine litter, in relation to the stage of decomposition and the carcase quality of the marine mammal.

**Table 15-1.** Guidelines for tissue sampling considering carcass DCC. Shading: green V indicates the process is of potential use in carcasses of the indicated DCC; grey (V)indicates that there may be limitations and red V indicates the procedure is not recommended/very unreliable, due to post mortem autolysis (IJsseldijk et al., 2019).

Analytical procedure	D C C 1	D C C 2	D C C 3	D C C 4	D C C 5	Comments/recommendations
Genetics	~	~	~	~	~	For DCC4 or 5: paleopathological procedures may be required on account of degraded DNA (eg extracting DNA from bone medulia)
Diet and marine debris	~	~	~	~	(✓)	If GIT is not intact, eg from post mortem scavenger damage, results are compromised
Age determination	~	~	~	~	(✓)	
Fatty acids and stable isotopes	~	~	~	~	(✓)	Depending on analysis planned
Parasitology	~	~	~	~	(✓)	Depending on analysis planned
Morphometrics	~	~	~	(✓)	(✓)	Girth measurements can be disrupted by bloating due to autolysis in DOC4-5
Gross pathology	~	~	~	(✓)	(✓)	Recommended for DOC4-5 in cases of forensic investigation
Reproductive studies	>	~	~	(√)	8	
Toxicology	>	~	~	(√)	8	Depending on pollutants. DCC1-2 for biomarker investigation.
Ear investigation	<	~	<	×	×	Inner ear analysis specifically: DCC1, histopathology of fixed ears possible up to DCC3
Microbiology	~	~	(√)	(√)	×	Depending on analysis planned. For DCC3-4 microbiology can still be worthwhile for detection of certain bacteria and fungi using specific culture methods. Should a septicaemia be suspected in DCC3-4 animals, then microbiological investigations should be undertaken on the kidney, as this is resilient to microbial post mortem invasion using specific culture methods.
Histopathology	~	~	(✓)	(√)	×	Recommended for DCC4-5 in cases of forensic investigation
Virology	~	~	(√)	×	×	Depending on analyses planned.

Biotoxins	~	~	(✓)	×	×	
Gas bubble analysis	~	~	×	8	×	If this procedure is conducted: it should be done first, before undertaking further assessments and dissections, particularly prior opening any part of the vascular system or removing the head.
Serology	~	(√)	(√)	8	×	Advisable both on blood serum and on cerebro-spinal fluid, the latter of which should be collected as soon as possible. In heavily autolyzed specimens, alternatives are "juice" obtained from skeletal muscle or lung, vitreous humour or pericardial fluid
Clinical chemistry	~	×	×	8	×	Vitreous humour is a possible option in decomposed cases . Care is needed however to ensure sufficient baseline data are available for the analyte in the species under investigation.

# Discovery circumstances – including entanglement and bycatch

Note the **circumstances** among the 5 categories:

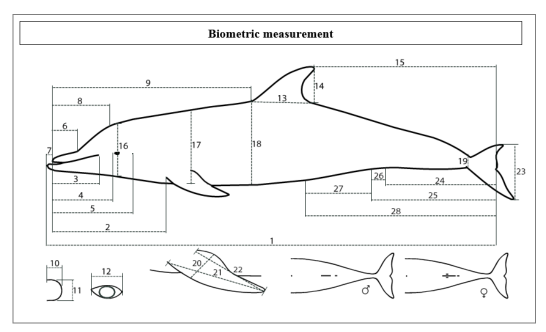
- Stranding\*: Animal found stranded on the beach or in the shoreline,
- *By-catch\*/Fisheries*: Animal accidentally captured by fishers (e.g. ingestion of a hook, trapped in a net, brought back by fishers, etc.) during fishing operations.
- Found at sea: Animal discovered on sea surface.

\* If possible, the <u>type of interaction with human activities and impact observed or suspected on dead</u> <u>or live stranded individuals</u> should be deduced from external or organs observations during the necropsy and complemented with veterinarian examinations. Also, an inspection of the oral cavity should be conducted for the presence of foreign material.

# Biometric Measurements and sex determination

Several basic and optional body lengths can be measured (in centimetres, precision 0.01 cm), as well as the weight.

The sex (male or female) should be noted, which can be determined by observation of sexual characters. Otherwise, specify by NI (for Not Identified).



*Figure 15-3.* Cetacean biometric measurement (from http://mammiferimarini.unipv.it/).

# Necroscopy and Health status

Necroscopy should be performed under the authorization needed at National level and with the presence of a veterinarian.

The body condition of a cetacean can be assessed by looking along the dorsal axis of the animal (poor, fair, good). The dorsal muscle mass (epaxial muscle) to either side of the dorsal fin of a robust animal will be rounded or convex. A thin animal will have a slight loss in epaxial muscle girth and could have a minor sunken aspect to the dorsal-lateral body. An emaciated animal will have a greater loss of epaxial muscle girth and will be concave down the dorsal-lateral body. Emaciated animal may also have more prominent indentation at the nape.

In addition visual inspection of the animal's fat reserves at the dorsal fin is recommended. Choose among the 3 categories:

- Thin;
- Fat;
- Normal.
- Not recorded (NR)

# Extraction of the gastrointestinal system:

- Expose the gastrointestinal system (GI) by removing all excess attached tissues, the heart and liver of the animal. Clamp the oesophagus proximal to the mouth and clamp the colon, the closest to the anal orifice.
- Remove the entire GI and place it on the examination surface or isolate the different portions of GI (oesophagus, stomach, intestines) by strangling and cutting between 2 clamps the gastro-oesophageal sphincter and the pyloric sphincter. This operation is easier if done by at least 2 operators.
- During the whole procedure, airborne contamination should be prevented as much as possible.

# Gut content analysis and marine litter isolation

- Before opening up the digestive tube, examine the outer wall to observe possible perforations by foreign bodies or areas of necrosis. Also, note any eventual secondary lesions, particularly a peritonitis following on a perforation of the digestive tube, an invagination of the digestive tube, an occlusion, etc. Photograph every lesion observed, taking care to get an overall view as well as close-up (macro-lens) photographs. Pictures must be stored referring to the code corresponding to the animal examined, describing the lesion in the description of the subject.
- The three parts of the gastrointestinal system (i.e. oesophagus, stomach, intestines) should be removed by adding a second strangling at the cut edge to prevent spillage of the contents. Each GI section should be opened lengthways using a scissor and slide the material directly out of the section onto a 1mm mesh sieve. The content should be cleaned with abundant tap water to remove the liquid portion, the mucus and the digested unidentifiable matter. Content should be inspected for the presence of any tar, oil, or particularly fragile material, and should be subsequently removed and treated separately. It should be then reported in the column "Notes" of the sampling sheet. All the material should be rinsed, collected in the 1mm sieve, and should be placed in tubes or in zipped bags, reporting the sample code (individual code, respective GI section) and stored at -20 °C, pending the laboratory analyses.
- NOTE: At this stage, for the optional differentiation of litter and microlitter, the material should be slid out of the section directly onto a 5mm mesh sieve superposed on a 1mm mesh sieve. Then, proceed with the rinsing and the storing of the material collected as described above, for both 1- and 5-mm sieves, reporting the samples code (individual code, respective GI section and size class (>5mm or 1-5mm)).
- If possible, follow the protocol developed in Corazzola et al (2021), which allows the simultaneous multidisciplinary analysis of GI by the implementation and standardization of a new methodological approach to the GIT of marine mammals. This protocol allows the collection of samples for different disciplines at the same time, performing the respective analyses, interpret and compare their results in a multidisciplinary way. The compatibility of multiple analyses allows the gaining of more information about the cause of death of stranded marine mammals and to enhance the knowledge of their biology and ecology.

The limit detection for MSFD is 1 mm. Building on the findings of the testing phase of the Plastic Busters MPAs project, it's recommended to also examine the fraction 0.1-1 mm.



*Figure 15-4.* A new prototype to isolate macro and microplastics in the gastrointestinal tract of stranded cetaceans (Corazzola et al., 2021). (Photo ©UNISI).

# Tissue collection

Before sampling the contents of the GI for the subsequent contaminant analysis, collect about 10g of each of the following tissues (level 1-4), wrap them in aluminium paper and store at -20 ° C:

- Muscle
- Liver
- Blubber (include skin) fat taken at the base of dorsal fin
- Kidney
- Brain (if possible include cerebrum and cerebellum)

In case of Level 1 specimen (max 1-2h after death):

- Blubber (include skin) for analysis of biomarkers analysis and contaminant analysis: take 10-20g from preserved in aluminum paper, store in liquid nitrogen or dry ice, and then place at -80 °C.
- Liver for biomarkers analysis and contaminant analysis: 10g in aluminum paper, store in liquid nitrogen or dry ice, and then place at -80 °C.Blood for contaminant analysis: 5-10 ml in tubes and store store in liquid nitrogen or dry ice, and then place at -80 °C.Each tissue stored in aluminium foil or Eppendorf must be labelled with the standard identification code of the animal.

# 15.3. Protocol for free-ranging marine mammals



Figure 15-5. Skin biopsies: a nonlethal tool for monitoring cetaceans. (Photo ©UNISI).

# **Cetaceans: Remote dart biopsy sampling procedure**

A number of successful studies show that cetacean skin biopsies are a powerful nonlethal tool for assessing ecotoxicologic risk in marine mammals and aspects of feeding ecology and food preferences.

Biopsy samples can be taken between the dorsal fin and the upper part of the caudal peduncle upon approaching the animal at a suitable distance and speed as specifically permitted for the species and

research project. The skin biopsy needs to be stored immediately in the proper conditions required for intended analyses. Common storage conditions include frozen, as is, in liquid nitrogen, dry ice, and at  $-80^{\circ}$ C and  $-20^{\circ}$ C freezers for long-term storage or stored either cold or at room temperature in cell medium, buffer, or specific reagents (e.g. RNA later). Skin biopsy is a powerful tool for ecotoxicologic studies for the following reasons: (1) it allows collection of a large number of samples across a wide geographic range; (2) it allows collection of sequential samples from the same animal if identified by photo identification or genetics; (3) it is suitable for residue analysis of many contaminants s; (4) it is suitable for several biomarker analyses and cell and organotypic cultures.

# Sampling procedure

- Skin biopsies (epidermis and dermis/blubber) from free-ranging dolphins (such as *Tursiops truncatus, Stenella coeruleoalba*) can be obtained using an aluminium pole armed with biopsy tips (e.g. 0.7 cm Ø, 3.0 cm length) or with a crossbow and darts.
- Skin biopsies from large odontocete (*Physeter macrocephalus*) or mysticete species (such as *Balaenoptera physalus* or other baleen whales) can be obtained with a crossbow and darts armed with tips (e.g. 0.9 cm Ø, 4.0 cm length).





*Figure 15-5.* Skin biopsy collection close to the dorsal fin of a fin whale and a striped dolphin. (Photo ©UNISI).

This type of sampling requires special permits issued by the competent authorities at national level.

# Skin biopsy collection

Once the biopsy has been collected from the animal, it should be processed as soon as possible.

1. Unscrew the tip from the arrow using gloves, put the biopsy in a small bag and write on the bag the code of the animal (put it in the fridge or keep as cold as possible). If there is more than one animal to collect at the same time and you cannot process the biopsy immediately, use a refrigerated bag until the processing.



*Figure 15-6.* Arrow and tip (left) and tip on the aluminum pole (right) with the collected biopsy. (Photo ©UNISI).

2. Remove the biopsy from the tip using tweezers, paying attention to keep the biopsy entire and put the biopsy on a clean petri dish.

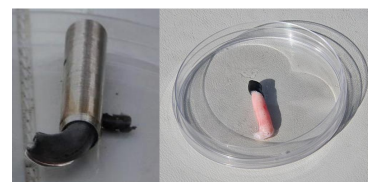
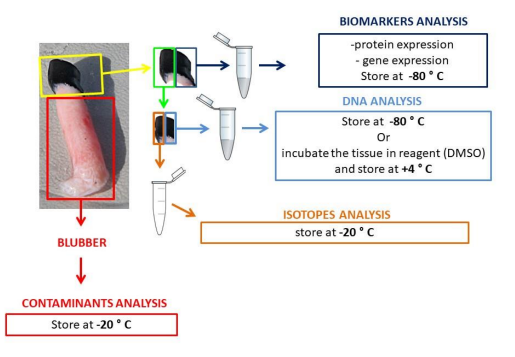


Figure 15-7. Biopsy sampled: left) biopsy tip, right) biopsy on a Petri dish. (Photo ©UNISI).

3. With a clean scalpel cut at least two pieces of skin (about 0.2x0.2 cm each) from the top of the biopsy (yellow squares) and put the separated aliquots of skin in two 0.5 ml Eppendorf. Whenever possible, for larger biopsy, divide the sampled biopsy in 4-5 different aliquots. Wrap up the skin+blubber (red square) in a small aluminium foil and put the biopsy in a 2 ml

Eppendorf (contaminants and protein analysis). Organotypic slice cultures should be potentially performed in specialized laboratory.

Write with a marker the code of the animal on the Eppendorf tubes, and, if possible, put inside the 2 ml tube a small piece of paper with the code of the biopsy written with the pencil, in order to be sure to not lose the name of the sample. During the operation, fill in the sampling sheet.



*Figure 15-8.* Biopsy with blubber (red square) and dermal part (yellow squares). Operational procedures.

- 4. Place the tubes in liquid nitrogen. The samples stored in RNAlater can be kept at room temperature for 24 hours and then stored at +4 °C or -20 °C for long-term storage.
- 5. Clean accurately the tips and boil them in freshwater for ten minutes to avoid cross contamination and pathogen transmission among individuals. If boiling the tips is not possible, rinse them with ethanol. Rinse with ethanol also the scalpel and the tweezers.

# Storage conditions

For skin biopsy:

- Dermal tissues (skin):
  - 40-60 mg in cryo-vial frozen at -80 °C (protein expression analysis/-omics analysis)
  - 30-50 mg in RNAlater at -20 °C or in cryo-vial frozen directly at -80 °C (gene expression analysis/transcriptomics)
  - 20-30 mg in 20% saturated DMSO with NaCl or in cryo-vial frozen directly at -80 °C (sex determination and genetic analysis)
  - 20-30 mg in cryo-vial frozen directly at -80 °C (stable isotopes analysis)
- Blubber tissues (fat):

• entire blubber in aluminium foil directly at -80 °C (contaminants analysis)

# Faeces collection

- For free-ranging cetaceans, faeces collection can be occasional and discontinuous, and generally, available only for fin whale. In case of localization of faeces, they should be collected as much as possible with a net (mesh size 200 μm or less) and put in falcon tubes for subsequent analysis: liquid nitrogen for contaminants and biomarker analysis, -20 °C for litter analysis.
- For monk seals, faeces should be entirely collected following the protocol by Lusher and Hernandez-Milian (2018) and stored at -20 °C or dry ice/liquid nitrogen for subsequent litter analysis, contaminant analysis and biomarker analysis. Food remains should be stored in 70% alcohol for diet analysis.

# **15.4.** The threefold approach in marine mammals

After the sampling phases described above (both in stranded and free-ranging animals), the analytical phases can be proceeded, following the methodologies applied in the testing phase of the Plastic Busters MPAs project.

The application of the **threefold approach** can elucidate not only the rate of ingestion in cetaceans, but also the multiple sub-lethal stresses that marine litter ingestion can cause in the short and long term. Each of the three investigation tools that make up the threefold approach can be applied independently or simultaneously using different methods according to the species and whether the animal is stranded or free-ranging.

The threefold approach comprises the following elements:

- Analysis of gastrointestinal content: For stranded cetaceans, it is possible to detect the occurrence and rate of marine litter ingestion and any associated pathology through analysis of the gastrointestinal content, with a particular focus on plastics and microplastics.
- Analysis of the levels of plastic additives, as a proxy for ingestion: An indirect approach can be used for free-ranging as well as stranded animals. The levels of plastic additives and associated PBT compounds can be measured to evaluate the exposure to marine plastic pollution.
- Analysis of biological end-points: Biomarker responses and omics analysis can be used to detect the potential toxicologic effect related to PBT and plastic additives related toplastic ingestion in free-ranging individuals or in stranded organisms up to a few hours after death.

# 15.5. Litter analysis and classification

# Macrolitter detection in stranded organisms

- Sort prey or litter items from the gastrointestinal tract into separate categories under a stereomicroscope, taking care of recording their weight.
- Measure the size of litter items and classify litter.

In addition, the following parameters should be recorded:

• For all categories (litter and other elements): the dry mass (grams, precision 0.01 g) of each category; dry the sample at room temperature during 24h minimum or in a stove at 35°C during 12h.

- For litter categories only: the number of fragments and items in each category: a fragment is a piece of litter that can be identified, whilean item is a set of fragments that seem to originate from the same piece of litter
- For the plastic litter categories only the total number of plastic fragments per colour category, with specifics as follow:
  - Total number of white-transparent plastic fragments;
  - Total number of dark coloured plastic fragments (black, blue, dark green...);
  - Total number of light coloured plastic fragments (cream, yellow, pink, light green...).
- Analyse at least 10% of the detected plastic by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

# Microlitter detection

- Examine the filter in the Petri dish under a stereomicroscope for particles resembling microplastics. Cover the filter with glass lids during observation to avoid the contamination of the sample.
- Photograph, count and record the type, colour and maximum length of microplastic particles using image analysis software and categorize microplastic particles.
- Analyse at least 10% of the detected microplastics by FTIR (Fourier Transform Infrared Spectroscopy) or Raman spectroscopy to determine the polymer composition and confirm the polymer origin of the detected particles.

# Microlitter detection in faeces

For free-ranging faeces, samples should be dried and digested using KOH 10%, then the solution filtered and litter should be classified. The dry mass (grams, precision 0.01 g) of each category should be recorded after drying at room temperature for at least 24h or at 35 °C for 12 h.

# Litter categories

Categorize marine litter according to the categories showed in Table 15-1. The categorization of the gastrointestinal tract contents and excreta is based on the general "morphs" of plastics (sheet-like, thread-like, foamed, fragment, other) or other general rubbish or litter characteristics. This is because in most cases, particles can't be unambiguously linked to particular objects. But where is possible, under notes in datasheets, the items should be described and assigned a litter category number using the "Joint List" developed by the TSG ML group (Fleet et al., 2021). In addition, it is important to measure and quantify also natural items (food and/or no food).

**Table 15-1.** Classification of Marine Litter items plus Food remain and Natural no food remain (fromINDICIT 2018).

BIOTA categories for contents of digestive tract				
PLA	PLASTIC	acronym	all plastic or synthetic items: note number of particles and dry mass for each category	
IND	pellets	ind	industrial plastic granules (usually cylindrical but also oval spherical or cubical shapes exist)	
	probab ind?	pind	suspected industrial, used for tiny spheres (glassy, milky,) (= microbeads)	
	sheet	she	remains of sheet, eg from bags, cling-foil, agricultural sheets, rubbish bags etc	
USE	thread	thr	threadlike materials, eg pieces of nylon wire, net-fragments, woven clothing; includes 'balls' of compacted material	
	foam	foam	all foamed plastics, polystyrene foam, foamed soft rubber (as in matrass filling), PUR used in construction etc	
	fragments	frag	fragments, broken pieces of thicker type plastics, can be bit flexible, but not like sheetlike materials	
	other	Poth	any other, incl elastics, dense rubber, cigarette-filters, balloon-pieces, softairgun bullets, objects etc. DESCRIBE!!	
RUB	OTHER RUBBISH	acronym	any other nonsynthetic consumer wastes: note number of particles and (in principle) dry mass for each category	
	paper	рар	newspaper, packaging, cardboard, includes multilayered material (eg Tetrapack pieces) and aluminium foil	
RUB	kitchenfood	kit	human food remains (galley wastes) like onion, beans, chickenbones, bacon, seeds of tomatoes, grapes, peppers, melon etc	
NUD	other rubbish	rubvar	other various rubbish, like processed wood, pieces of metal, metal air-gun bullets; leadshot, paintchips. DESCRIBE	
	FISHHOOK	hook	fishing hook remains (NOT FOR HOOKS ON WHICH LONGLINE VICTIMS WERE CAUGHT - THOSE UNDER NOTES)	
POL	POLLUTANTS (INDUS/CHEM WASTE)	acronym	other non-synthetic industrial or shipping wastes (number of items and mass per category (wet for paraffin)	
	slag/coal	slag	industrial oven slags (looks like non-natural pumice) or coal remains	
POL	oil/tar	tar	lumps of oil or tar (also note as n=1 and g=0.0001g if other particles smeared with tar but cannot be sampled separately)	
	paraf/chem	chem	lumps or soft mush of unclear paraffin, wax like substances (NOT stomach oil!); if needed estimate mass by subsampling	
	featherlump	confea	lump of feathers from excessive preening of fouled feathers (n=1 with drymass) (NOT for few normal own feathers)	
FOO	NATURAL FOOD	foo	various categories, depends on the species studied, and aims of study	
NFO	NATURAL NON FOOD	nfo	anything natural, but which cannot be considered as normal nutritious FOOD for the individual	

# Collection of data

For each organism, an assessment is made of:

- 1. Frequency of occurrence (%) of ingested macro and microlitter for each species, calculated as the percentage of the individuals examined with ingested macro- and microplastics.
- 2. Abundance (N) of macro and microlitter ingested per individual (average number of items/individual) for each species, calculated as a total and per category. Since currently there are inconsistencies in the literature in reporting abundance of ingested litter, it is recommended to report average number of items per individual considering both all individuals examined and only individuals found with ingested macro and litter.
- 3. Total dry weight (g) of the detected waste expressed on grams (precision: second decimal place). This weight refers to each single category found in a specific organ (or faeces) of the specimen.

Other information as colour of items, polymer of the different items (at least 10% of the total items) and different incidence of litter in oesophagus, stomach and intestine, incidence and abundance are useful for research and impact analysis.

# 15.6. Analysis of plastic tracers and PBTs

# Plastic additives

The compounds to be detected are:

- <u>*Phthalates:*</u> a group of chemicals widely used as additives to make plastics more flexible and harder to break; they can interfere with endocrine system (Baini et al., 2018).
- <u>Bisphenol A</u>: used in the production of polycarbonate, can have endocrine disrupting effects (Crain et al., 2007; Halden, 2010; Oehlmann et al., 2009) and the styrene and polyvinyl chloride monomer, used in the production of polystyrene and polyvinyl chloride (PVC), can be carcinogenic and/or mutagenic (Lithner et al., 2011; Papaleo et al., 2011; Xu et al., 2004).
- <u>Polybrominated diphenyl ethers:</u> they belong to the group of brominated flame retardants (BFRs), which are used in various polymeric materials such as plastic parts, resins, textiles, and other substrates to reduce their fire hazards (BSEF 2003; Król et al. 2012).

# Persistent, bioaccumulative and toxic substances (PBTs)

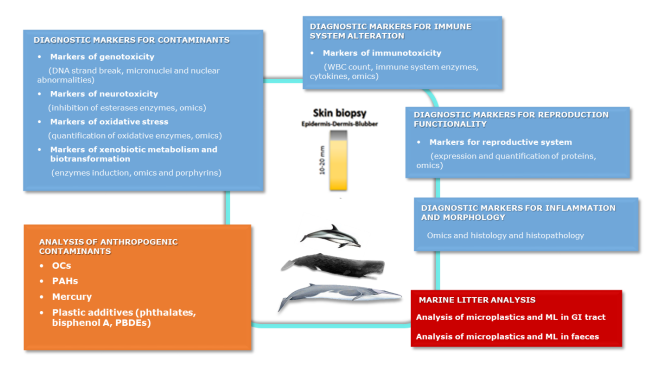
In addition to the plastic additives that may leach from plastics when released into the marine environment, plastics also tend to adsorb in their surface persistent bioaccumulative and toxic substances (PBTs) (e.g. organochlorine compounds OCs, PAHs and PBDEs) and metals (e.g., lead, copper and cadmium) that are present in the seawater.

Depending on the compounds and the tissue to be analysed, different methods should be applied to detect the presence of plastic related contaminants in the sentinel species (Annex VI).

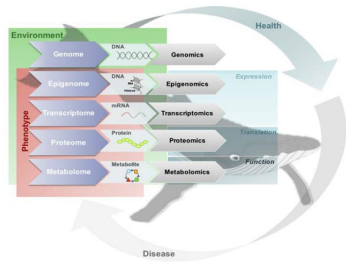
# **15.7.** Biomarkers analysis

The toxicological effects associated with the presence of marine litter can be evaluated using a set of diagnostic and prognostic methodologies, by means of biomarkers. A non-exhaustive list of existing biomarker approaches and plastic tracers' contaminants that are usually applied in marine mammal analyses is reported in Annex VI.

Biomarkers have been selected on the basis of the level of biological responses and in relation to the main effects related to marine litter/microplastics ingestion. The selected biomarkers can diagnose different impacts related to: a) physical damages/effects of marine litter, b) exposure to/effect of chemical tracers, and c) exposure to/effect of adsorbed chemicals.





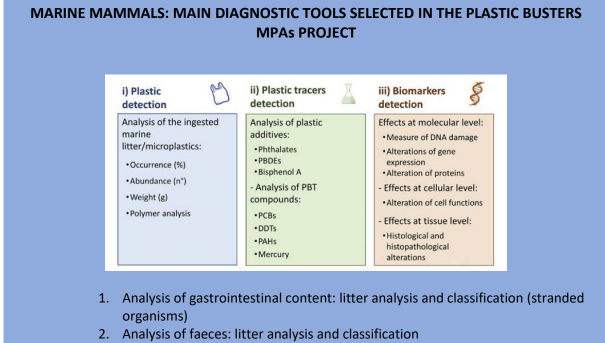


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**Figure 15-8.** A) three-fold approach to detect the marine litter presence and impacts to marine mammals. B) Omics techniques in skin biopsies (from Mancia 2018).

Starting from this initial list and building on the findings of the testing phase of the Plastic Busters MPAs project, the most suitable diagnostic tools to detect the presence and impact of ML on marine mammals are proposed here below.

**Table 15-2.** Main diagnostic tools selected in the PLASTIC BUSTERS MPAs project to detect the presence and impact of ML in marine mammals.



- 3. Analysis of plastic tracers: phthalates
- 4. Analysis of biological end-points: gene expression (*adipoq, ahr, gr, ppara, pparg, thra, thrb, cd36, cyp1a, cyp3a*), Omics

# 15.8. Materials & Equipment for sampling

The following material and equipment are necessary for the correct application of the protocol (stranded organisms):

- Boots
- Camera
- Clamps (at least 6) or roast wire
- Clips with claws
- Containers for samples (Bottle/zipped bags)
- Cooler
- Cut-resistant gloves
- Garbage bag
- Glasses and protective mask or shield
- Nitrile Gloves
- Integral protective suit
- Measuring cylinders (2 L, 1L, 50cL; precision 0.1L)

- Measuring decimetre
- Measuring tape
- Metal containers
- Metal spoon
- Observation sheet
- Pen
- Permanent marker
- Precision balance
- Rope (to mark-off the zone)
- Sampling sheets
- Scalpel
- Scissors
- Sieve with 1 mm mesh
- Sieve with 5 mm mesh
- Transport bins or containers

The following material and equipment are necessary for the correct application of the protocol (free-ranging organisms):

- Aluminium foil
- Aluminium Pole
- Bicoculars
- Camera
- Crossobow
- Cryoboxes
- Cryovials
- Darts
- DMSO (20% saturated with NaCl)
- Eppendorf (0.5 ml. 1.5 ml. 2.0 ml)
- Ethanol (70%, 100%)
- Falcon tubes
- Glass Petri dishes
- Gloves
- GPS
- Liquid nitrogen dewar (in alternate dry ice)
- Net (for faces collection)
- Paper and block-notes
- Paper towels
- Pasteurs
- Pencils
- Permanent markers
- Plastic Sealable bags
- RNAlater
- Ruler
- Scalpels
- Spare batteries
- Spare camera batteries and memories
- Thermic bags

- Tips (for crossbow and aluminium pole)
- Tweezers
- VHF Radio

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# 15.9. Sampling & recording sheets

# Monitoring Marine Litter (Macro-Micro) in biota: stranded marine mammals

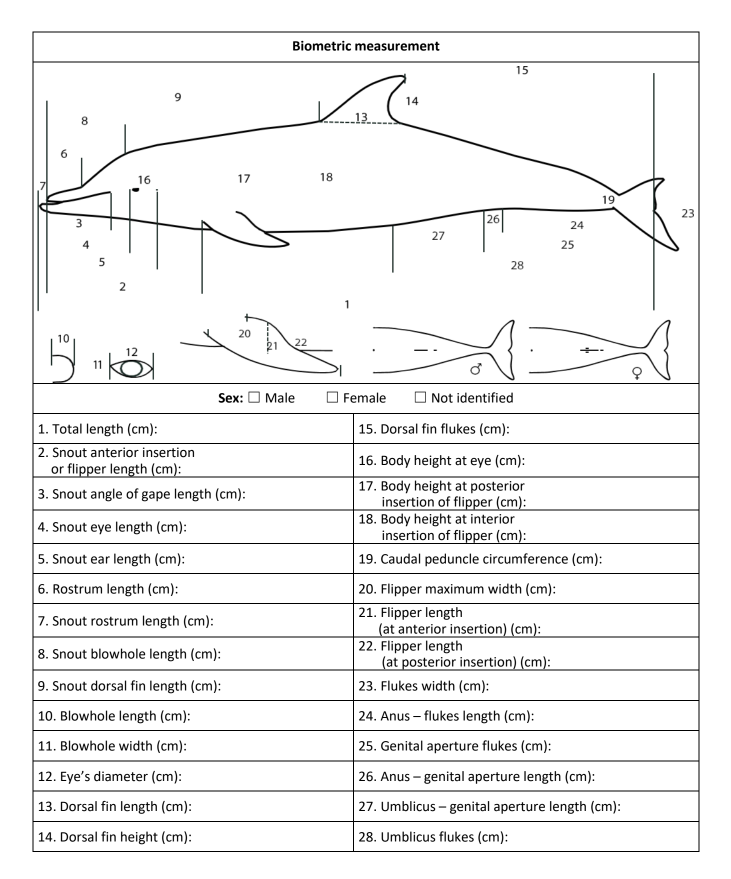
Species:		ID code:			
Location/Country:		Latitude	Longitude		
Discovery size metonoos	□ By catch/Fishery	□ Found at sea	□ Stranding		
Discovery circumstances	Dead at rescue cer	iter 🗌 Other			
<b>Discovery circumstances</b> (Please specify according to the toolkit)					
Date of discovery					
Date of necroscopy					

Animal body condition								
Conservation status	🗆 Level 1	🗌 Level 2	🗆 Level	3	🗆 Level 4			
Health status	Poor	🗌 Fair	🗆 Good	I				
Main iniurias	No injuries	🗌 Fractu	re 🗌 Amp	utation	Sectioning			
Main injuries	□ Abrasion							
Affected parts	□ Wings	□Tail	□Neck	🗌 Hea	d 🗆			
Affected parts	Other							
Fat reserve	🗆 Thin	🗆 Fat	🗆 Normal	🗆 No	t recorded (NR)			

	Collected tissues	N°. of aliquots
Muscle		
GI tract		
Liver		
Fat tissue		

Necropsy performed by:

Name and Institution:



# Notes and remarks:

# Monitoring Marine Litter (Macro-Micro) in biota: free-ranging cetaceans

Species:			ID code:					
Sampling date and time:								
Sampling site:			Latitude		Longitude			
Sampling tool:							N	
□ Single	□ Couple □	□ Group	p Dimension of the group:					
Picture ref.:								
Weather cond	Weather condition Sea:			Wind:				
Lenght (m):								
Aprox. age:	Δ	🗆 Sub-adult			🗆 Juvenile			
Side sample and position:  Right  Left								
	Contaminants (blubber)	Liquid N <sub>2</sub>	RNA later	DMSC	)	Cell medium	Bouin	
	PCR (skin)							
Aliquots	WB (skin)							
	Sex (skin)							
	lsotopes (skin)							
	Omics							
	Histology							
	Treatment Dose		Time	Hour-notes				
Slices								
Faeces								

Notes and remarks:

## **ANNEX VI**

	CHEMICAL COMPOUND	TISSUE/SAMPLE	ANALYTICAL METHOD
PLASTIC ADDITIVES	Phthalates	Blubber, muscle, liver, whole organism, skin biopsy	Baini et al., (2017), Fossi et al., (2016), Savoca et al., (2018), Routti et al., (2021)
		Muscle	Ballesteros-Gómez et al., (2009)
	Bisphenol A	Blubber, skin biopsy	Xue et al., (2016)
		Blood	Cobellis et al., (2009)
	Polybrominated diphenyl ethers	Blubber, muscle, liver, blood, skin biopsy	Muñoz-Arnanz et al., (2016), (Zaccaroni et al., (2018), Bartalini et al., (2019), Baini et al., (2020), Aznar-Alemany et al., (2021)
ADSORBED CONTAMINANTS	Polycyclic aromatic hydrocarbons	Blubber, muscle, liver, blood, skin biopsy	Marsili et al., (2001)
	Organochlorine contaminants	Blubber, muscle, liver, blood, skin biopsy	Marsili and Focardi, (1997), Bartalini et al., (2019), (Genov et al., 2019), Baini et al., (2020), Aznar-Alemany et al., (2021)
	Mercury	Blood, skin, skin biopsy	Correa et al., (2013)

 Table A-9. Tissues and methods to be used to detect plastic tracers in marine mammals.

**Table A-10.** Biological end point detected in free-ranging marine mammals by the biomarker and Omics approach. The analysis on blood, liver, kidney and muscle can be performed only in dead marine mammals (level 1 - fresh carcass).

EFFECT	TISSUE	TEST
GENOTOXICITY	Blood, skin	Comet assay (Molino et al., 2019) (*) Mn test (Bolognesi et al., 2006) ENA assay (Casini et al., 2018); (Pacheco and Santos, 1997)
OXIDATIVE STRESS	Liver, kidney	CAT, GST, LPO, GPX, GR, GSH (Yu et al., 2018) (*)
UNIDATIVE STRESS	Plasma, skin	LPO (Fossi et al., 2016), Casini et al., 2018) CAT (Fossi et al., 2013)
ΙΜΜυΝΟΤΟΧΙCITY	Blood	Total and differential white blood cells (WBC) count (Casal and Orós, 2007; Davis et al., 2008; Caliani et al., 2019) H:L ratio (Caliani et al., 2019) Respiratory burst (Secombes, 1990; Caliani et al., 2019) TAS assay (Miller et al., 1993; Caliani et al., 2019) Lisozyme enzyme (Keller et al., 2006; Caliani et al., 2019) casp8, casp9, TRAF (Karami et al. 2017; Mathieu-Denoncourt et al., 2015) (*)
PERPOPULATION	Plasma, Gonads	CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015) (*) Vitellogenin (Fossi et al., 2004)
REPRODUCTION	Plasma, skin	Vitellogenin (Herbst et al., 2003) CYP17A, CYP19, ERs, VTG, StAR (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*)
HISTOPATHOLOGY INFLAMMATION AND MORPHOLOGY	Liver, kidney	Histopathology, histology (Pedà et al. 2016; Karami et al. 2017; Batel et al., 2018) (*)
XENOBIOTIC METABOLISM AND BIOTRANSFORMATION	Blood, skin, faeces, liver	CYP1A; AHR, CYP3A (Fossi et al. 2014, Panti et al. 2011; Rochman et al., 2013) (*) Porphyrins (Guerranti et al., 2014) (*)
NEUROTOXICITY	Muscle	AChE, BChE (Barboza et al., 2018) (*)

CELLULAR STRESS	Blood, skin, liver, kidney	PPARA, PPARG, HSP70, GPX, E2F1 (Mathieu-Denoncourt et al., 2015; Panti et al., 2011) (*) Gamma glutamyl transferase (GGT) (Nematdoost Haghi and Banaee, 2017) (*) Cortisol and corticosterone (Flower et al., 2015) LDH (Nematdoost Haghi and Banaee, 2017) (*)
OMICS	skin	Epigenetics (Mancia et al., 2021), Transcriptomics (Lunardi et al., 2016)

(\*) effects detected after laboratory or field exposure with MPs or Plastic Tracers.

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