Genetic Fact Sheet

Dab Scophthalmus rhombus







PANDORA

Paradigm for Novel Dynamic Oceanic Resource Assessments

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Genetic Fact Sheets

Review of available genetic information on population structuring in exploited species

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PANDORA Project

The Blue Growth of European fisheries is at risk due to over-exploitation, unforeseen changes in stock productivity, loss of markets for capture fisheries due to aquaculture, future trade agreements opening European markets to external fleets, and fluctuations in the price of oil and other business costs. All of these risks need to be considered when providing advice needed to sustainably maximize profits for the diverse array of fisheries operating in European waters and to help safeguard the benefits this sector provides to the social coherence of local, coastal communities.

PANDORA aims to:

1. Create more realistic assessments and projections of changes in fisheries resources (30 stocks) by utilising new biological knowledge (spatial patterns, environmental drivers, food-web interactions and density-dependence) including, for the first time, proprietary data sampled by pelagic fishers.

2. Advise on how to secure long-term sustainability of EU fish stocks (maximum sustainable/"pretty good" and economic yields) and elucidate tradeoffs between profitability and number of jobs in their (mixed demersal, mixed pelagic and single species) fisheries fleets. Provide recommendations on how to stabilize the long-term profitability of European fisheries.

3. Develop a public, internet-based resource tool box (PANDORAs Box of Tools), including assessment modelling and stock projections code, economic models, and region- and species-specific decision support tools; increase ownership and contribution opportunities of the industry to the fish stock assessment process through involvement in data sampling and training in data collection, processing and ecosystem-based fisheries management.

The project will create new knowledge (via industry-led collection, laboratory and field work, and theoretical simulations), new collaborative networks (industry, scientists and advisory bodies) and new mechanisms (training courses and management tools) to ensure relevance, utility and impact.

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List of abbreviations

GFCM	General Fisheries Commission for the Mediterranean
ICES	International, Council for the Exploration of the Sea
IUCN	International Union for Conservation of Nature
IUU	Illegal, Unreported and Unregulated
MSY	Maximum sustainable yield
SAC	Scientific Advisory Committee
STECF	Scientific, Technical and Economic Committee for Fisheries
SD	Subdivision
TAC	Total Allowable Catch

North-East Atlantic ICES subareas, divisions and subdivisions

- 1) Subarea 1 Barents Sea
- 2) Subarea 2 Norwegian Sea, Spitzbergen and Bear Island
- 3) Subarea 3
 - Division 3.a, Skagerrak (subdivision 20) and Kattegat (subdivision 21)
 - Division 3.b-c, Sound (subdivision 23) and Belt Sea (subdivision 22)
 - Division 3.d, Baltic Sea (subdivisions 24-32)
- 4) Subarea 4 North Sea (divisions 4.a-c)
- 5) Subarea 5 Iceland (division 5.a) and Faroes Grounds (division 5.b)
- 6) Subarea 6 West of Scotland (division 6.a) and Rockall (division 6.b)
- 7) Subarea 7
 - Irish Sea (division 7.a), West of Ireland (division 7.b), Porcupine Bank (division 7.c)
 - Eastern English Channel (division 7.d), Western English Channel (division 7.e)
 - Bristol Channel (division 7.f), Celtic Sea (divisions 7.g-h), Southwest of Ireland (divisions 27.7.j-k)
- 8) Subarea 8
 - North and Central Bay of Biscay (divisions 8.a-b)
 - South Bay of Biscay (division 8.c)
 - Offshore Bay of Biscay (division 8.d), West of Bay of Biscay (division 8.e)
- 9) Subarea 9 (Portoguese Waters)
- 10) Subarea 10
 - Azores Grounds (division 10.a) and Northeast Atlantic South (division 10.b)
- 11) Subarea 11 (incorporated in FAO Fishing Area 34)
- 12) Subarea 12 North of Azores
 - souther mid-Atlantic Ridge (division 12.a)
- 13) Subarea 13 (incorporated in FAO Fishing Area 34)
- 14) Subarea 14 East Greenland, Northeast Greenland (14.a), Southeast Greenland (14.b)



How to read the factsheets

Genetic structure factsheets are presented for each species. Current knowledge on genetic population structure is summarised and compared with stock units used in assessment and management. The presence of mismatches is emphasised as well as priorities for future work. At the beginning of the factsheets, a summary is presented with green-yellow-red color symbols for 'Population structure', 'Match between genetic and stock assessment units' (units for which scientific advisory bodies, as ICES and the GFCM, provide advice on stock status and fishing opportunities), 'Match between genetic and management units' (units for which TACs are set by the European Council), 'Match between stock assessment and management units'. The information in the factsheet is organized in the following sections:

Distribution: general information can be found on the distributional range of the species, with a focus on the NE Atlantic, Mediterranean and Black Sea.

Current management status: an overview is provided on the current management and assessment units present for the species in European Seas. The importance of the species for each fishery is included, reporting if the species is mainly a by-catch or if direct fishery exists for the stocks. A mismatch between stock assessment and management units already exists for certain species and it is showed in Table 2.

Genetic population structure in a nutshell: provides the key take-home messages, both in terms of current knowledge on genetic population structure and in terms of priorities for future work. In this section, an overall picture of population structure of the species is given, based on considerations on the type of markers, sampling designs and findings of the included studies. It is also discussed if genetic evidence supports the stock assessment and management units currently in use.

Mismatch: in this section the mismatch between genetic and stock assessment/ management units is highlighted. Two types of mismatch can be observed. Here, we refer to '*Type I*' mismatch when a genetically homogeneous population is assessed/managed in multiple stock units (oversplitting); while we refer to '*Type II*' mismatch when genetically different populations are wrongly considered part of the same stock assessment/management unit (undersplitting).

Summary of genetic evidence: in this section a more detailed summary of the studies is provided in a chronological way. In general, the type of genetic markers used by different studies depends on the widely available markers at the time. Early studies used allozymes and often reported a lack of differentiation among sample locations. However, later studies using the more highly polymorphic microsatellites and SNPs showed presence of differentiation even in areas where it was not previously detected. Conversely, in other cases presence of differentiation was reported at few allozyme loci, not confirmed



subsequently with strictly neutral markers. This and other contradictions between studies were addressed if possible. Advances in sequencing technology, as well as the use of more sophisticated statistical analysis and sampling design to maximise the detection of population structure have made enormous changes in the awareness we have of genetic structure in marine fish species (Hauser and Carvalho, 2008). Most of the mismatches found in initial studies between genetic population structure and stock assessment and management units were due to a lack of differentiation reported between samples assessed/managed in different units (referred to as 'Type I' mismatch in Table 1). However, these mismatches are often solved by more recent investigations, that applied highly polymorphic markers, as well as a sampling design that maximise the chance of detecting population structure, i.e. collecting individuals in spawning aggregations. Particular emphasis should be placed on the sampling season and individuals included in the analysis that are extremely important factors for the detection of population structure in marine fish species (Nielsen et al. 2009b). Moreover, despite in previous studies a neutral background of low differentiation was commonly detected, recently the application of markers under selection allowed the detection of high levels of differentiation and occurrence of locally adapted populations. Therefore, a summary of genetic studies found in literature is provided. For each study, sampling design, temporal and spatial analyses and markers used have been critically evaluated. Strengths and shortcomings of the available studies are reported and based on these considerations an overview is given.

Table 2.1. Summary table of available information on genetic population structure and match between genetic, assessment and management units of commercial fish species exploited in the NE Atlantic, Mediterranean and Black Sea.

Species	No. Studies	Population structure	Match genetic- Stock assessm ent units	Match genetic- managem ent units	Match stock assessment -manage- ment units	IUCN status
Dab, Limanda limanda	2	yes	yes	-	-	LC

IUCN Abbreviations: NE= Not evaluated, DD= Data Deficient, LC= Least Concern, NT= Near Threatened, VU= Vulnerable, EN= Endangered, CR= Critically Endengered. Eu= Europe, Glo= Global, Med= Mediterranean (IUCN 2021).



FACT SHEET

Dab, *Limanda limanda*

Number of studies Population structure

Match genetic- Stock assessment units

Match genetic- Management units

Match Stock assessment- Management units



Distribution¹

Dab, *Limanda limanda* (Linnaeus, 1758), is a demersal flatfish species commonly found in the North-East (NE) Atlantic shelf from the Bay of Biscay to Norway, as well as in the White Sea, Barents Sea, Baltic Sea and Iceland. Dab is a very common species in the North Sea (Daan et al. 1990).

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Current management status

There are two ICES stocks in the NE Atlantic for dab (Figure 3.3): one in the North Sea, Skagerrak, Kattegat and the other in the Baltic Sea. Dab in mainly a by-catch species in the direct fishery of cod and in mixed fisheries of other flatfish (plaice and flounder) of higher commercial importance. Discard has been estimated to be close to 50% for the Baltic stock (ICES 2020b). In the Baltic Sea, it is present only in the western part of the basin (Subdivisions 22-25), in fact the majority of landings is reported from the Belt Sea (Subdivision 22), with smaller amount from subdivision 24 and 25, while the occurrence of individuals in the eastern part (Subdivisions 26-32) is rare (ICES 2020b). Total landing in

the Baltic Sea for 2019 is 1102 t, mostly fished by Germany and Denmark in mixed fisheries of flatfish species (ICES 2020b). Due to a lack of studies, dab inhabiting the North Sea, the Skagerrak and Kattegat was considered as a single stock assessment unit (ICES 2016). Fish are mainly caught as by-catch in the demersal fishery for other flatfish species, mainly alongside plaice and sole (ICES 2018a). Currently, there are no TACs set for dab and ICES is not providing

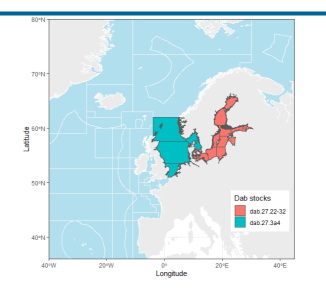


Figure 3.3. Dab ICES stock assessment units

¹ Further details on symbols and how to read the factsheet are provided on page 16



advices for fishing opportunities for Baltic and North Sea dab, but only information on the status of the Baltic Sea stock (ICES 2020b).

Genetic population structure in a nutshell

Microsatellites and SNPs have been used to analyse population structure of dab around the British Isles (Tysklind et al. 2013) and in the North Sea, Baltic Sea transition zone (Le Moan et al. 2019a), respectively. Genetic evidence supports the existence of separate populations of dab in the North Sea, Irish Sea and Baltic Sea. Presence of population admixture and hybridization between North Sea and Baltic Sea individuals in the transition zone was reported.

Mismatch

In contrast with the current stock assessment units for dab, the presence in the Kattegat and the transition zone of individuals of admixed origins was reported. Further investigations are needed to define the boundaries of the North Sea dab stock. Further analysis should also focus on the transition zone to unravel the extent of hybridization and population admixture of the North Sea and Baltic Sea stocks.

Summary of genetic evidence

Since Reiss's review (2009), in which an absence of genetic information on population structure for dab was reported, two studies have been published (Table 1). Tysklind et al. (2013) using microsatellites showed the presence of two temporally stable populations inhabiting the North Sea and the Irish Sea. The importance of considering population structure of species that, similar to dab, are used as bioindicators was highlighted. In fact, the biomarker responses of dab in UK waters may be population specific (Tysklind et al. 2013).

Genetic population structure of dab in the North Sea, Baltic Sea and the transition zone (the Kattegat, the Belt Sea and the Øresund) was analysed by SNP markers by Le Moan et al. (2019a). The presence of two populations and a continuum of hybridization along the transition zone, with substantial population admixture, was reported. The divergence between North Sea and Baltic Sea populations was supported (F_{ST} = 0.020) (Le Moan et al. 2019a). Genetic divergence was associated with environmental gradients of salinity, sea surface temperature and sea bottom temperature in the transition zone.



Table 1. Summary table of genetic population structure studies of commercial marine fish species exploited in the North-East Atlantic Ocean, Mediterranean and Black Sea.

Species	Region	Sampling locations	No. Samples (Number of individuals)	Spawning	Maturity	Life stage	Genetic Marker	Differentiati	Mismatch genetic-SA	Mismatch genetic- MZ	LA, LG, MSA	Reference
Dab	NE Atlantic	NS (4), EC (2), IS (7), Atlantic (2)	39 (3006) *	у	у	Ad	Msat (14) ^N	Yes	no	na		(Tysklind et al. 2013)
	NE Atlantic	NS (1), NBTZ (3), BAL (2)	6 (148)	у	na	na	SNPs (3468) ^s	Yes	Type II	na	LG, LA	(Le Moan et al. 2019a)

Table 2. Mismatch between stock assessment (SA) units and genetic population structure (Type I and II explained) and mismatch between management and genetic units.

Species	Stock assessment unit	Mismatch SA unit - genetics (Type II)	Mismatch SA unit - genetics (Type I)	Management units	Mismatch management unit - genetics
Dab,	<u>dab.27.22-32</u>		Hybridization and		
			population admixture in		
Limanda	dab.27.22-32		the NS-BS transition zone		
limanda			(Le Moan et al., 2019)		



The following abbreviations are used for the geographic locations: North-East Atlantic (NE Atlantic), Mediterranean Sea (Med), Northwest Atlantic (NWA), Adriatic Sea (Adr), Aegean Sea (Aeg), Africa (AFR), Alboran Sea (Alb), Atlantic (Atl), Atlantic Iberian (Atl IB), Australia (AU), Azores (Azo), Baltic Sea (BAL), Barents Sea (BS), Bay of Biscay (BOB), Black Sea (BLS), British Isles (BI), Canada (CAN), Canary (Cn), Cantabrian Sea (Cant), Celtic Sea (CS), English Channel (EC), Faraday Seamount (Far), Faroe Islands (FRO), fjord (fj), Galicia (Gal), Greece (GRC), Greenland (GRL), Gulf of Cadiz (GC), Gulf of Lion (GoL), Hebrides (Heb), Iceland (ICE), Ionian Sea (Ion), Ireland (IRE), Irish Sea (IS), Irminger Sea (Irm), Kattegat (Kat), Lake Mogilnoe (Mog)Lofoten (Lof), Madeira (Mad), Marmara Sea (MS), Mid-Atlantic Ridge (MAR), Morocco(MOR), Namibia (Nam), New Zeland (NZL), Newfoundland and Labrador (NL), North Sea (NS), North Sea-Baltic Sea Transition zone (NBTZ), Norway (NOR), Nova Scotia (Nov), Porcupine Bank (Por), Portugal (PRT), Reykjanes Ridge (Reyk) , Rockall Bank (Roc), Russia (RUS), Scotian Shelf (SS), Scotland (SCO), Shetland (SHE), Sicily (SIC), Skagerrak (Ska), Spain (SPA), Svalbard and Jan Mayen (SJM), Tasman Sea (TS), Tunisia (TUN), Tyrrhenian Sea (Tyr), White Sea (WS); north (n), south (s), east (e), west (w), central (c); Norwegian Coastal Cod (NCC), North-East Arctic Cod (NEAC).

For each study the species, sampling locations (for abbreviations see below) and in brackets the number of samples are shown; the total number of samples and individuals analysed is reported, as well as the number of temporal replicates in superscript or (*) if multiple temporal replicates are included. The spawning, maturity and life-stage of samples included are summarised as follow, *Spawning*: y= if samples collected in spawning season/grounds are included, na= not available, no= samples outside spawning season/grounds. *Maturity*: y= mature individuals included; na= maturity not available; no= immature individuals. *Life-stage*: Ad= adult; juv= juveniles; lar= larvae; eg= eggs; na= not available. Genetic markers (All= allozymes; Msat= microsatellites; Minisat= minisatellites; SNPs= Single Nucleotide Polymorphisms; mtDNA= mitochondrial DNA; Cyt-b= cytochrome b; COI= Cytochrome c Oxidase subunit I; COIII= Cytochrome c Oxidase subunit III; CR= Control Region; RAPD= Random Amplified Polymorphic DNA); number of loci or base pairs analysed in brackets, in superscript S= if at least one locus is under selection, N= neutral markers (only if neutrality was tested). Differentiation, if genetic differentiation was detected (Yes, No). Mismatch genetic- SA= mismatch of the genetic units found and the stock assessment units. Mismatch genetic- MU = mismatch of genetic units with the management units. We refer to *'Type II'* mismatch when a genetically homogeneous population is assessed/managed in multiple stock units (oversplitting); while we refer to *'Type II'* mismatch when genetically different populations are wrongly considered part of the same stock assessment/management unit (undersplitting). LA= Local Adaptation, LG= Landscape Genetics, MSA= Mixed Stock Analysis.