

Non-indigenous seaweeds in the Northeast Atlantic Ocean, the Mediterranean Sea and Macaronesia: a critical synthesis of diversity, spatial and temporal patterns

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Non-indigenous seaweeds in the Northeast

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Abstract

- 55 Effective monitoring and combatting the effect of non-indigenous seaweeds relies on a solid
- 56 confirmation of the non-indigenous status of the species. We critically analysed the status of
- 57 presumed non-indigenous seaweed species reported from the Mediterranean Sea, the Northeast
- 58 Atlantic Ocean and Macaronesia, resulting in a list of 140 species whose non-indigenous nature is
- 59 undisputed. For an additional 87 species it is unclear if they are native or non-indigenous
- 60 (cryptogenic species) or their identity requires confirmation (data deficient species). We discuss the
- 61 factors underlying both taxonomic and biogeographic uncertainties and outline recommendations to
- 62 reduce uncertainty about the non-indigenous status of seaweeds. Our dataset consisted of over

- 63 19,000 distribution records, half of which can be attributed to only five species (Sargassum
- 64 muticum, Bonnemaisonia hamifera, Asparagopsis armata, Caulerpa cylindracea and Colpomenia
- 65 peregrina), while 56 species (40%) are recorded no more than once or twice. In addition, our
- analyses revealed considerable variation in the diversity of non-indigenous species between the
- 67 geographic regions. The Eastern Mediterranean Sea is home to the largest fraction of non-
- 68 indigenous seaweed species, the majority of which have a Red Sea or Indo-Pacific origin and have
- 69 entered the Mediterranean Sea mostly via the Suez Canal. Non-indigenous seaweeds with native
- 70 ranges situated in the Northwest Pacific make up a large fraction of the total in the Western
- 71 Mediterranean Sea, Lusitania and Northern Europe, followed by non-indigenous species with a
- 72 presumed Australasian origin. Uncertainty remains, however, regarding the native range of a
- 73 substantial fraction of non-indigenous seaweeds in the study area. In so far as analyses of first
- 74 detections can serve as a proxy for the introduction rate of non-indigenous seaweeds, these do not
- 75 reveal a decrease in the introduction rate, indicating that the current measures and policies are
- 76 insufficient to battle the introduction and spread of non-indigenous species in the study area.

77 Highlights

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- Non-indigenous seaweed species in the Northeast Atlantic Ocean, the Mediterranean Sea and Macaronesia are critically reanalysed.
- >19,000 distribution records revealed considerable variation in diversity of non-indigenous
 seaweed species in the study area.
- Taxonomic and biogeographic uncertainties hamper a critical evaluation of the nonindigenous status of many seaweed species.
- 84 **Keywords:** biodiversity, Chlorophyta, Europe, invasive alien species, non-indigenous species,
- 85 Phaeophyceae, Rhodophyta

Introduction

- 87 Over the course of several centuries, human-mediated transport has led to the introduction and
- 88 establishment of more than 14,000 non-indigenous species in Europe (EASIN, 2022). Some of
- 89 these non-indigenous species profoundly affect the abundance, diversity, interactions and evolution
- 90 of native biota and consequently affect ecosystem structure, functions and services (Simberloff et
- 91 al., 2013; Dawson et al., 2017; Blakeslee et al., 2020). The introduction of non-indigenous species
- 92 can also result in substantial negative economic impacts (Hulme et al., 2009). The reported costs of

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biological invasions, at a global level, were estimated to be at least 1.288 trillion US Dollars over 1970–2017 (Diagne et al., 2021). Furthermore, biotic homogenisation and consequently also the impact of non-indigenous species on native ecosystems are expected to increase in the context of climate change (Bennett et al., 2021). The management of biological invasions depends heavily on lists of reliably identified nonindigenous species. Such lists form an essential tool underpinning prevention, control, mitigation or eradication strategies (Kolar & Lodge, 2001), and in particular to facilitate prevention and early detection, which are the most cost-effective for management (Simberloff et al., 2013). In addition, government and management agencies use lists of non-indigenous species in their policies to protect nature and reverse the degradation of ecosystems. For instance, the primary criterion for the descriptor D2 dedicated to non-indigenous species under the European "Marine Strategy Framework Directive" is the rate of novel introductions per 6-year period (European Commission et al., 2021). Comprehensive and accurate lists of non-indigenous species, their respective origin, and geographical and temporal spread are therefore crucial for an effective response and legislation to battle threats imposed by non-indigenous species. Unfortunately, the compilation of such lists is marred by the challenges involved (McGeoch et al., 2012; Costello et al., 2021). At local scales, lists may be confounded by limited occurrence data and hence underestimate the number and spread of non-indigenous species. At a more fundamental level, taxonomic uncertainty and the associated lack of expertise in species identification are regarded as severe problems (Zenetos et al., 2017). The effects of taxonomic uncertainty are likely more pronounced for less studied taxa and poorly sampled regions. For example, upon re-examination of about 100 potential non-indigenous taxa of marine molluscs, almost half of the records turned out to be misidentifications or the distributional data were incorrect (Zenetos et al., 2017). While DNA-assisted identification has the potential to solve identification problems, misidentifications of entries in genetic databases combined with geographic and taxonomic sampling bias make it a challenge in itself to correctly interpret gene sequence data (Viard et al., 2019; Fort et al., 2021; Tran et al., 2022). In addition, taxonomic knowledge is not static. Evolving taxonomic insights, often derived from genetic and biogeographic studies, alter our views on the indigenous or non-indigenous nature of taxa, requiring checklists to be continuously updated (Taylor, 2010; Guareschi & Wood, 2019). This problem is exacerbated in the marine environment where many cryptic species have been documented (Appeltans et al., 2012). The above-mentioned problems related to lists of non-indigenous species definitely apply to seaweeds, which represent one of the largest groups of marine non-indigenous organisms, constituting between 20 and 29% of all marine non-indigenous species in the Northeast Atlantic

127 Ocean, the Mediterranean Sea and Macaronesia (hereafter referred to as "the study area") (Schaffelke et al., 2006; Molnar et al., 2008; Katsanevakis et al., 2013) (Fig. 1; Fig. 2). The 128 129 consequences of non-indigenous species on native ecosystems have only been studied in a very limited number of species. Although some non-indigenous species have been observed to have 130 positive ecosystem effects (e.g. Gracilaria vermiculophylla in the Venice Lagoon and Northeast 131 Atlantic mudflats; Davoult et al., 2017; Sfriso, 2020), impact studies on such seaweeds have mostly 132 133 detected negative ecological effects, with reduction in abundance of native biota being most 134 frequently reported (Williams & Smith, 2007; Weinberger et al., 2008; Hammann et al., 2013; Katsanevakis et al., 2014; Maggi et al., 2015; Bulleri et al., 2017; Anton et al., 2019). However, 135 136 contrary to the evidence of substantial negative impact on coastal ecosystems of many non-137 indigenous seaweeds (e.g. Caulerpa cylindracea, Caulerpa taxifolia, Codium fragile), so far Rugulopteryx okamurae is the only seaweed included in the list of invasive alien species of Union 138 concern (COMMISSION IMPLEMENTING REGULATION (EU) 2022/1203 of 12 July 2022 139 140 amending Implementing Regulation (EU) 2016/1141). This EU regulation enforces member states to adopt measures to prevent, minimise or mitigate the adverse impact of those species. 141 Regional lists of non-indigenous seaweed species have been regularly published until 142 143 recently. For the Mediterranean Sea, which has been disproportionately affected by non-indigenous species as well as other stressors (Lejeusne et al., 2010; Katsanevakis et al., 2014), non-indigenous 144 seaweeds have been critically revised on a regular basis (Verlaque et al., 2015; Zenetos et al., 2017; 145 146 Galil et al., 2021). Non-indigenous seaweeds of Macaronesia were included in Borges et al. (2010), 147 Chainho et al. (2015), Gallardo et al. (2016) and Castro et al. (2022). Bárbara et al. (2005) and 148 Brodie et al. (2016) provided a list of non-indigenous seaweeds as part of a revised check-list of Galician and British seaweeds, respectively. However, there are gaps and uncertainties for some 149 150 regions, and more importantly, a critical compilation encompassing the Northeast Atlantic Ocean, Macaronesian and Mediterranean regions is currently lacking. The absence of a critically revised 151 152 list in the study area not only impedes a comprehensive overview of non-indigenous seaweeds, but 153 may also introduce ambiguity related to the status of specific taxa due to differences in the criteria used to define non-indigenous species (see Materials and Methods). In addition, in the absence of a 154 comprehensive list, spatial and temporal patterns of introductions are difficult to deduce. 155 156 To address this knowledge gap, we compiled a database of non-indigenous seaweeds in the Northeast Atlantic Ocean, the Mediterranean Sea and Macaronesia with their distribution records, 157 158 their likely origin and putative introduction vectors. These data are used to provide a quantitative assessment of the spatio-temporal dynamics of primary and secondary introductions and to detect 159

shortcomings in the monitoring and legislation required to tackle the introduction of non-indigenous species more effectively.

Materials and methods

consensus (AlgaeBase, Guiry & Guiry, 2023).

Data compilation

We compiled a database of non-indigenous marine seaweed species records from three regions, namely the Northeast Atlantic Ocean (excluding Greenland), the Mediterranean Sea and Macaronesia (Fig. 2). For some of the analyses we subdivided the Northeast Atlantic Ocean into Lusitania and Northern Europe and the Mediterranean Sea into a Western and Eastern part. With respect to Macaronesia, the compilation includes records from the Azores, Canary Islands, Madeira and the Salvagen Islands, but not Cape Verde. The dataset builds on previous lists by Mineur et al. (2010) and Verlaque et al. (2015), and includes published records of species occurring in a natural environment and flagged as non-indigenous in the study area irrespective of taxonomic confidence and biogeographic status (see below). In addition, we included unpublished records produced by various research projects conducted by, amongst others, the Station Biologique de Roscoff (France), National Biodiversity Data Centre (Ireland), Stichting ANEMOON (the Netherlands), Scottish Natural Heritage (Scotland), the ICES Working Group on Introductions and Transfers of Marine Organisms 2004, as well as the European Alien Species Information Network (EASIN, 2022) records, collection data, GBIF records and personal data. All records were added to the database

The species listed as non-indigenous include those that are naturalised (i.e. having established permanent, self-maintaining populations), as well as species for which no information is available on population status (i.e. species referred to as 'alien' by Verlaque *et al.* 2015). Species that have been demonstrated to be misidentifications or unsupported records are excluded from the list. To promote consistency in definitions and criteria used to determine whether a species is non-indigenous, we have adopted the criteria for assessing the biogeographic status proposed by Essl *et al.* (2018) (Fig. 3). This framework stresses 1) the need for crossing a biogeographic barrier, 2) the involvement of direct or indirect human agencies in the physical movement of individuals, spores or fragments, and 3) the ability of the species to reproduce without human assistance in the introduced range. The combination of these criteria excludes records of species which are in the process of

under the name they were reported as. Names were updated according to the most recent taxonomic

expanding their range naturally, for example as a result of global warming. Species entering the Mediterranean Sea through the Suez Canal (i.e. Lessepsian migrants), on the other hand are considered non-indigenous because of the anthropogenic nature of the dispersal corridor. In contrary, species entering the Mediterranean Sea through the strait of Gibraltar, without a human-vector, are not considered as non-indigenous. The dataset also includes species indigenous to the study area that have demonstrably become displaced within he study area as a result of human-mediated exchanges. Examples include exchanges of species between Atlantic and Mediterranean shores. Species for which the area of origin is unknown are assigned as 'cryptogenic' (sensu Carlton, 1996). In cases where there is not sufficient information to be conclusive on their biogeographic status, species are labelled as 'data deficient'. Species with low uncertainty, for which there is no doubt about their non-indigenous status, have been labelled 'non-indigenous'.

Added to these criteria but highly relevant with respect to seaweeds, where a solid taxonomic framework is often lacking for many taxa, is taxonomic confidence. We assigned a 'high-confidence score' (score = 1) to accepted nominal species that had not been shown to be a species complex based on molecular studies in their introduced or native ranges. A high score was also assigned to species for which, so far, there is sufficient confidence in unambiguous identification based on morphology. We acknowledge, however, that the latter does not rule out the potential existence of cryptic species hidden under the accepted nominal species. Conversely, species that belong to an understudied complex of cryptic species were assigned a low-confidence score (score = 0). A cryptic species is defined here as a taxon composed of two or more species that have been classified as a single nominal species, because they were initially not distinguished based on their morphological characteristics (Bickford *et al.*, 2007; Pante *et al.*, 2015).

Recognising we cannot be conclusive about the non-indigenous status of many seaweed species, we explicitly acknowledge the uncertainty in the assessment of the taxonomic as well as biogeographic status of putative non-indigenous seaweeds in the study area (Fig. 3). The status of each species is concisely described in Suppl. Material Table S1.

For every species we determined the year when the species was first reported in the Northeast Atlantic Ocean, Mediterranean Sea and/or Macaronesia. Where possible, this date refers to the year the species was detected (i.e. collection date) rather than when the record was published (i.e. publication date). We acknowledge that detection dates may not portray the actual date the species was introduced. For each species an estimate is provided for its native biogeographic range. If the native range could not be assessed, we indicated 'uncertain'. The putative distribution of the

species was based on literature reports included in AlgaeBase (Guiry & Guiry, 2023). Species traits (e.g. thallus size) were obtained from AlgaeTraits (Vranken *et al.*, 2022). For spatial and temporal analyses, distribution records were filtered on a combination of unique year, coordinates and species name to eliminate potential duplicate records. The complete dataset has been archived at Zenodo and is available at DOI: 10.5281/zenodo.7798640. This dataset contains the following information for each record: currently accepted scientific name, the scientific name under which it was originally reported, year of record, location, country, coordinates and reference.

Results and Discussion

A total of 19,724 records of non-indigenous seaweeds were collected dating from 1808 to 2022 (Fig. 2). Of these, 17,104 were retained after removing duplicates and incomplete data. The geographical distribution of the records highlights considerable sampling of non-indigenous seaweeds from all coastlines in the study area (Fig. 2). The list contains 227 species (Table 1). Non-indigenous species make up approximately 10% of the seaweed flora in the Mediterranean Sea, 6% in the Northeast Atlantic Ocean and 4% of the Macaronesian flora. The total number of 227 includes all species regardless of taxonomic and biogeographic uncertainty. For 84 species neither their non-indigenous status nor their taxonomy is challenged (Fig. 4). These species make up 83% of the distribution records in the database. Half of the distribution records can be attributed to only five species (Sargassum muticum, Bonnemaisonia hamifera, Asparagopsis armata, Caulerpa cylindracea and Colpomenia peregrina). Fifty-six species are most likely non-indigenous, but decisions are hampered by taxonomic uncertainties (Fig. 4). On the other hand, 87 of the 227 species have a cryptogenic or data deficient status (30 species with an uncertain biogeographic status, and 57 species for which both the geographic status and taxonomic confidence are uncertain), meaning that the evidence for a non-indigenous status is mediocre to weak (Fig. 4).

We emphasise that the categorisation of putative non-indigenous species according to biogeographic uncertainty and taxonomic confidence emerged as a consensus among the authors of this paper. A literature search will undoubtedly reveal several additional species names that could potentially be added to the list of cryptogenic or data deficient species. However, there is little added value in incorporating species names which are wholly unsupported or most likely result from misidentifications or other mistakes. Evidently, both taxonomic and biogeographic uncertainty plague the compilation of lists and databases of non-indigenous species. Below we discuss how the

level of sophistication of systematic and biogeographic knowledge translates to uncertainty in the number of non-indigenous seaweeds in the study area.

Taxonomic confidence

For 140 species the non-indigenous nature of the species itself is not disputed. However, the reliable identification of 56 of those species is challenging, and therefore their current distribution as well as their putative region of origin are questionable. In most cases this uncertainty can be attributed to a poorly established taxonomic framework. Taxonomic uncertainty is rife in seaweeds. In the absence of DNA sequence data the identification of many seaweed species is particularly difficult (e.g. Van Oppen *et al.*, 1996; Maggs *et al.*, 2007; Cianciola *et al.*, 2010; De Clerck *et al.*, 2013; Verbruggen, 2014). Taxonomic uncertainty is much higher among small-sized species: 67% of species smaller than 5 cm are flagged as taxonomically uncertain, compared to 34% of species larger than 5 cm. Of the taxa larger than 5 cm with high taxonomic uncertainty are many that belong to genera that are widespread in tropical and warm-temperate regions (e.g. *Avrainvillea, Caulerpa, Codium, Dichotomaria, Ganonema, Hypnea*). From a biogeographic perspective, taxonomic uncertainty plagues "only" 20% of species with a Northwest Pacific origin (11 of 56 species), but 62% of species with a likely Lessepsian or tropical Indo-Pacific origin (26 of 42 species) (Table 1).

Recent advances in the taxonomy of several genera, nearly always assisted by DNA sequence data, have demonstrated that many so-called wide-ranging (or cosmopolitan) seaweeds actually consist of species complexes of morphologically almost indistinguishable species (pseudocryptic species), or even truly cryptic species which are indistinguishable based on morphological criteria. The individual species are often confined to specific geographic areas (e.g. Won et al., 2009; Vieira et al., 2017; Diaz-Tapia et al., 2018; Leliaert et al., 2018; Diaz-Tapia et al., 2020). A more refined taxonomic framework therefore alters our understanding of the biogeography of the species in many cases and consequently our interpretation of their native versus non-indigenous status. The Caulerpa racemosa complex is highly representative of how evolving insights into species diversity alter our views of the taxa being non-indigenous in the study area. While initially Caulerpa specimens with vesiculate branchlets collected in the Mediterranean Sea were identified as C. racemosa, the latter proved to be a complex of at least eight species, three of which (C. chemnitzia, C. cylindracea and C. requienii) are currently considered non-indigenous in the Mediterranean Sea (Verlaque et al., 2000; Verlaque et al., 2003; Draisma et al., 2014; Verlaque et al., 2015). Similarly, a better understanding of the taxonomy of foliose Grateloupia species

resulted not only in the recognition that *G. turuturu* was introduced in the study area from the
Northwest Pacific, as opposed to *G. doryphora* whose distribution is likely to be restricted to the
Pacific coast of South America (Gavio & Fredericq, 2002), but also revealed that so-called nonindigenous foliose *Grateloupia* species in the study area were actually a mixture of two nonindigenous species, *G. lanceolata* and *G. turuturu*, and a native species, *G. lanceola*, which had
been regarded a synonym of *G. doryphora* (Verlaque *et al.*, 2005; Figueroa *et al.*, 2007).

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In many other instances, however, conspecificity of populations from the native and nonnative regions remains to be demonstrated. There are also examples where several non-indigenous
species are thought to be conspecific by some authors but regarded as distinct species by others. For
example, some authors consider *Antithamnion hubbsii* distinct from *A. nipponicum* (Athanasiadis,
1996), while others treat the former as a synonym of the latter (e.g. Kim & Lee, 2012). Similarly,
records of *Polysiphonia morrowii* and *P. senticulosa* likely belong to the same species, even though
both species are regarded as distinct (D'Archino *et al.*, 2013; Stegenga & Karremans, 2015;
Piñeiro-Corbeira *et al.*, 2020). Given the widespread nature of cryptic and pseudocryptic diversity
in seaweeds, continuous efforts of DNA-assisted identifications through Sanger sequencing will
probably continue to revise our view on non-indigenous species.

Although DNA sequence data are in many cases a great help in verifying species identities, this does not mean DNA solves every single problem like a magic wand. Apart from reference sequences in repositories not being available or reliable, patterns of genetic divergence can be complicated and prone to different interpretations. For example, differences in the interpretation of genetic patterns and species boundaries in the genus Melanothamnus led to the recognition of a single species, M. harveyi s.l. (McIvor et al., 2001) or by contrast to the recognition of at least three separate species, including M. akkeshiensis, M. japonicus and M. harveyi s.s. (Savoie & Saunders, 2015). The narrower species concept would result in an interpretation whereby M. harveyi is native to the Northeast Atlantic Ocean rather than a non-indigenous species introduced to the study area from the Northwest Pacific Ocean. Under the alternative scenario which recognises a single genetically diverse species, M. harveyi is widely distributed globally with both cryptogenic and non-indigenous haplotypes in the study area (Piñeiro-Corbeira et al., 2019). One should note that despite the availability of a good number of sequences of these species/haplotypes, the potential native area of the species (Northwest Pacific Ocean) has been scarcely sampled. Therefore, it is still possible that M. harveyi s.s. can be present in this region but remained undetected. Distribution records of M. harveyi and M. japonicus are included as M. harveyi/japonicus in our dataset.

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Biogeographic uncertainty A lack of baseline data with respect to the global distribution of seaweeds is the major contributor to biogeographic uncertainty reported for 87 taxa (Fig. 4). Brown and green seaweeds display slightly less biogeographic uncertainty, 28% and 34%, respectively, compared to 42% for red seaweeds. Baseline data of seaweed diversity along coastlines in the study area as well as the putative native regions in the form of herbarium collections, censuses and historical checklists can serve as a reference for the presence of species in a given area. Here as well, low confidence in the taxonomy and identification of seaweeds makes the interpretation of species lists exceedingly difficult. If a species is not reliably identified, its distribution is not reliable. As a result, biogeographic and taxonomic uncertainties usually go hand-in-hand. Three different categories of factors that lead to biogeographic uncertainty are discussed below. Pseudo-indigenous species Several seaweed species have been described from the study area that were presumed native, but later turned out to be non-indigenous species. Carlton (2009) named such species pseudoindigenous. For example, Dictyota cyanoloma was described as a new species from the Mediterranean Sea and Macaronesia (Tronholm et al., 2010), but subsequent collecting efforts revealed that the species most likely represents a cryptic introduction (Aragay Soler et al., 2016; Steen et al., 2017; Tran et al., 2021). Similarly, Porphyra olivii described from Greece (Brodie et al., 2007a) turned out to be conspecific with Neopyropia koreana (Vergés et al., 2013; Yang et al., 2020), a species native to the Northwest Pacific. Such insights invariably result from DNA-assisted species identification and subsequent interpretations of biogeographic patterns. Hereby widely disjunct distribution ranges are interpreted as non-natural and therefore the result of humanmediated dispersal. Evidently, determining the non-indigenous nature of a species becomes more difficult for historic introductions. In such cases we fully rely on DNA signatures which can point toward a nonindigenous nature of the species. DNA-assisted identification of historic voucher specimens of Codium fragile revealed that the invasive (sub)species was already introduced into the study area as early as 1845 (Provan et al., 2008), which is roughly a century before phycologists realised the species was actually native to the Northwest Pacific Ocean and non-indigenous to the study area as well as several other parts of the world. In the case of Cutleria multifida, described from Norfolk,

England as early as 1801, genetic signatures point toward an introduction of the Mediterranean Sea populations from the Northwest Pacific. The Northeast Atlantic Ocean populations, however, are genetically more diverse and well-differentiated from those in Japan and are therefore considered native (Kawai *et al.*, 2016). It remains to be determined if *Cutleria multifida* is native to the Northeast Atlantic Ocean as well as the Pacific, or whether an even more complex history of historic introductions underlies this pattern.

Discerning natural from human-mediated dispersal

Eventually, the possibility of introductions needs to be evaluated against historic and ongoing natural dispersal events. The recent observation of Flabellia petiolata from the south coast of England confronts researchers with exactly this question (Díaz-Tapia et al., 2020). Despite a long tradition of seaweed studies and regular surveys, F. petiolata was never recorded from the British Isles prior to 2013. The closest populations of the species are found in the Mediterranean Sea and the Canary Islands. A recent introduction would be the most obvious explanation. However, the English populations of F. petiolata could also be interpreted as a relic of a formerly more widespread Atlantic Ocean distribution. The species' range might have been continuous during warmer periods in the Holocene, but persisted in the Northeast Atlantic Ocean in a handful of refugia during colder periods. Afterall, several native species, e.g. Cladophora battersii, Codium bursa and Halopithys incurva display similar distribution patterns (Maggs & Hommersand, 1993; Brodie et al., 2007b).

Quaternary climatic cycling probably also facilitated dispersal of temperate species across the tropical Atlantic Ocean connecting southern Africa with Europe. The presence of *Schizymenia apoda* in the Azores, the British Isles and Namibia may have resulted from natural amphi-equatorial dispersal events in recent geological times, but also a human-mediated introduction in the Atlantic Ocean cannot be ruled out given the presence of *S. apoda* in Australia and China (Gabriel *et al.*, 2019; Gunnarsson *et al.*, 2020). Natural dispersal events from the Northeast Pacific Ocean to Northern Europe through the Bering Strait may be difficult to discern from introductions (Lindstrom, 2001; Bringloe & Saunders, 2019). For instance, *Schizymenia jonssonii*, a species recently described from Iceland, may have colonised the northern Atlantic Ocean naturally via the Bering Strait but it is equally possible the species is a relatively recent introduction (Gunnarsson *et al.*, 2020).

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409 410 Population-level sampling and the application of genetic markers with sufficient intraspecific resolution (e.g. fast evolving spacer regions, microsatellite markers or SNP data) have the potential to shed light on natural versus human-mediated dispersal events, and more generally to help in reconstructing introduction history (Viard & Comtet, 2015), but are rarely used in studies of putative seaweed introductions. Notable exceptions include the invasive history of *Fucus* species, *Sargassum muticum* and *Gracilaria vermiculophylla*.

Coyer et al. (2011) demonstrated a North Pacific origin of Fucus distichus followed by at least two separate colonisation events of the North Atlantic Ocean prior to the last glacial maximum, which makes this species native to Europe. However, the taxon, having a predominantly northern distribution, was accidentally introduced in the Oslofjord followed by further expansion in the Kattegat region as a result of an introduction event in the late 19th century (Coyer et al., 2002, as F. evanescens, currently regarded as a subspecies of F. distichus). Fucus serratus was also exported from mainland Europe to Atlantic North America, Iceland and the Faroes (Coyer et al., 2006; Brawley et al., 2009). Discharging of ballast stones in destination harbours is considered the prime source of introductions in Atlantic North America and Iceland in the 19th century. The F. serratus population in the Faroes is of more recent origin (late 20th century) and was most likely introduced from Iceland (Coyer et al., 2006). Many marine benthic organisms including seaweeds but also many invertebrates display amphi-Atlantic distribution (Haydar, 2012). For such disjunct distributions, distinguishing scenarios of post-glacial relicts or natural long-distance dispersal from human-assisted dispersal (and introduction) is challenging and most often requires a combination of life-history traits assessment and high resolution molecular markers. The power of genome-wide genetic variation was demonstrated for Sargassum muticum (Le Cam et al., 2020); whereas microsatellite markers failed to reveal any genetic variation in the invaded range of the species, a panel of single-nucleotide polymorphisms (SNPs) obtained from ddRAD sequencing confirmed a secondary introduction to the Northeast Atlantic Ocean from the Northeast Pacific Ocean, but also revealed two additional cryptic introductions to Europe. Similarly, Krueger-Hadfield et al. (2017) identified the areas in the native region that most likely contributed to the European invasions of the red alga Gracilaria vermiculophylla. Subsequent work used SNPs to refine the origins and understand evolution during invasion (Flanagan et al., 2021).

In Macaronesia, several confounding factors further complicate the interpretation of the nonindigenous nature of species. The geographical location of Macaronesia, bordering the tropical Atlantic Ocean, contributes significantly to this difficulty in interpretation. Several tropical and

subtropical taxa are, probably erroneously, attributed a pantropical distribution, which not only 411 contributes to high taxonomic uncertainty, but the latter also translates into biogeographic 412 413 uncertainty. In addition, it is not always evident to preclude natural dispersal to explain the presence 414 of particular species. *Halimeda incrassata*, a species that naturally occurs in the tropical western 415 Atlantic Ocean (Verbruggen et al., 2006), was reported from Porto Santo, Madeira, by Wirtz & Kaufmann (2005) and more recently from the Balearic Islands by Alós et al. (2016). Further 416 417 surveys indicated the species is also present in the Canary Islands (Sangil et al., 2018) and Azores 418 (Costa et al., 2017). Even though in the Mediterranean Sea the species displays typical invasive 419 behaviour, the involvement of human activities in its establishment is not clear. Another species from tropical western Atlantic Ocean, Caulerpa ashmeadii, was recently reported from Porto Santo, 420 421 Madeira, and may represent a similar case of natural range expansion across the Atlantic (Ribeiro et al., 2023). Amphi-Atlantic Ocean distributions have been confirmed using molecular markers for 422 423 several seaweed taxa (Cladophoropsis membranacea, Leliaert et al., 2009; Laurenciella marilzae, 424 Cassano et al., 2012; Laurencia catarinensis, Machin-Sanchez et al., 2012; Tronholm et al., 2012; Vertebrata foetidissima, Díaz-Tapia et al., 2013; Dictyota spp., Tronholm et al., 2013; Caulerpa 425 prolifera, Varela-Álvarez et al., 2015; Laminaria digitata, Neiva et al., 2020; Lobophora spp., 426 Vieira et al., 2020). In these examples presumed natural distribution ranges have not been 427 428 challenged. 429 Also of note is that Macaronesia covers a large geographic area, encompassing several 430 biologically diverse archipelagos. Several taxa that have long been reported from the Canary Islands 431 and Madeira and are considered native in those areas, have recently been reported from the Azores. 432 The geographic position of the Azores and the actual oceanographic current circulation in the North 433 Atlantic Ocean would not seem favourable for natural range expansions from the Canary Islands and Madeira. In some cases, initial reports from anthropogenic habitats, such as harbour 434 environments (e.g. Caulerpa webbiana), favour the hypothesis of an anthropogenic factor in the 435 436 range expansion, although evidence is lacking for several other species which are presumed non-437 indigenous (e.g. Halimeda incrassata, Xiphosiphonia pinnulata, Hypoglossum heterocystideum). 438 A final category of uncertainty in natural vs. human-mediated dispersal mechanisms concerns 439 those non-indigenous species which have been displaced within the study area. The biogeographic 440 history of the Mediterranean Sea biota is closely intertwined with the Northeast Atlantic Ocean to which it is connected by the narrow Strait of Gibraltar (Bianchi & Morri, 2003; Patarnello et al., 441 442 2007; Le Gall *et al.*, 2021), which results in a subset of species being shared between both regions.

However, several Northeast Atlantic Ocean species have been recently introduced into the 443 Mediterranean Sea, often in lagoons with extensive aquaculture facilities, e.g. Ascophyllum 444 nodosum, Chorda filum, Fucus spiralis and Grateloupia minima (Petrocelli et al., 2013). In some 445 cases however, patterns become more complex, for example, when native and non-indigenous 446 populations co-occur as is the case for *Chondria coerulescens*, *Vertebrata fucoides* and possibly 447 also Ganonema farinosum (Verlaque et al., 2015). It is worth remembering that many 448 449 Mediterranean Sea species naturally dispersed from the Atlantic Ocean after the Zanclean flood 450 which occurred after the Messinian salinity crisis about 5.33 myr (Blondel et al., 2010). 451 Species of unresolved origin 452 In several cases the non-indigenous nature of certain seaweeds remains unresolved. Some recently 453 described species in the study area, such as *Polysiphonia radiata* and *P. delicata*, are mainly known from marinas and are probably non-indigenous but their origin remains unknown because it is likely 454 that they remained undescribed in their native area (Díaz-Tapia et al., 2017). Morphological 455 similarity of putative non-indigenous species to native species can also complicate interpretation of 456 non-indigenous patterns as demonstrated by Anotrichium furcellatum. The latter was originally 457 458 described from Naples, but considered non-indigenous in the Northeast Atlantic Ocean. The Mediterranean Sea populations, however, may have been largely replaced by a cryptic introduction 459 of A. okamurae, originally from the Northwest Pacific Ocean (Verlague et al., 2015). The status of 460 A. furcellatum and A. okamurae has not yet been tested with molecular data. 461 Similar arguments could be made for species that are considered native in the study area. In 462 463 the case of Lobophora delicata, which is not considered as non-indigenous, a lack of baseline data makes it difficult to be conclusive on its status as a native species. As pointed out by Vieira et al. 464 465 (2019), the first records of *Lobophora* in the Mediterranean Sea date back to 1955 (Edelstein, 466 1960), which is surprising for a distinctive seaweed genus which can be easily found in many places growing at a depth of 0.5 m. In contrast, other genera of Dictyotales were invariably reported from 467 468 the Mediterranean Sea in the 18th or early 19th century. Has L. delicata been overlooked or does 469 the late discovery of the species correspond to a more recent introduction? Without proper baseline data, e.g. herbarium records, this is difficult to test, and if the species does not display typical 470 471 invasive behaviour its native status may simply never come into question. 472 A puzzling case is presented by several taxa with clear Indo-Pacific affinities which 473 appeared in the Mediterranean Sea prior to the opening of the Suez Canal in 1869, e.g.

Acanthophora nayadiformis, Asparagopsis taxiformis and Ganonema farinosum. For instance, Asparagopsis taxiformis was first described from Alexandria in the Mediterranean Sea as Fucus taxiformis Delile (1813), and thus reported as a native species. However, further molecular work revealed that this accepted species was made of five distinct lineages, possibly corresponding to two cryptic species (Ní Chualáin et al., 2004; Andreakis et al., 2007; Dijoux et al., 2014), one of them presumably present in the Mediterranean Sea prior to the opening of the Suez Canal, and one more recently introduced. Similar complexity was revealed for the closely related species Asparagopsis armata, supposedly introduced in the study area, for which novel sampling in the South Pacific Ocean showed the existence of two highly divergent clades, presumably corresponding to two cryptic species, one of them distributed in Europe, South Africa and Tasmania, and one restricted (so far) to Western Australia, New Zealand and Tasmania (Dijoux et al., 2014). Such cases highlight the difficulty in establishing whether a species is non-indigenous in the absence of large sampling encompassing the global distribution of the targeted presumably non-indigenous species.

The spatial patterns and origins of non-indigenous seaweeds

Analysis of the distribution of non-indigenous seaweeds in the study area reveals clear patterns in richness and the number of species shared among regions. The large-scale spatial patterns are discussed below in a context of dispersal vectors that determine spread and establishment of non-indigenous species.

The Eastern Mediterranean Sea is home to the highest number of non-indigenous seaweeds (77 species), followed by the Western Mediterranean Sea (47 species) and Lusitania (45 species). Macaronesia and Northern Europe harbour somewhat lower numbers (36 and 40 species, respectively). These numbers refer to high-confidence non-indigenous species only. Adding cryptogenic and data deficient species further underscores the higher number of non-indigenous species in the Eastern Mediterranean Sea. In the latter region an extra 47 species are flagged as cryptogenic or data deficient, which is considerably higher compared to the other regions which typically host 20 or less cryptogenic and data deficient species. In all regions, most of the non-indigenous seaweeds belong to Rhodophyta (between 63-76% of the species non-indigenous in each region), followed by brown seaweeds (18-22%), while green seaweeds contribute to 8-18% of the species recorded (Fig. 5).

We did not detect a significant correlation between the number of non-indigenous species and the number of records (Pearson correlation = -0.18, p-value = 0.74), which indicates that differences of non-indigenous species between regions are not a mere artefact of sampling effort. The number of non-indigenous species also does not scale with the length of the coastline (Pearson correlation = 0.10, p-value = 0.88). As will be argued below, the number of non-indigenous species in a given region and the fraction of species shared between regions is a complex function including the efficiency of primary and secondary dispersal vectors combined with abiotic (and potentially biotic) ecological factors that determine the establishment of non-indigenous species in the recipient ecosystems (reviewed in Maitner *et al.*, 2021).

The Mediterranean Sea and the Northeast Atlantic Ocean share 45 high-confidence nonindigenous species, while Macaronesia shares roughly an equal number of non-indigenous species with the Mediterranean Sea (24 species) and the Northeast Atlantic Ocean (21 species) (Fig. 6A). Within the Northeast Atlantic Ocean, Macaronesia and Lusitania share 22 high-confidence non-indigenous species, while Macaronesia and Northern Europe share none other than the 14 non-indigenous species present in all three Northeast Atlantic regions (Fig. 6B). A relatively low number, 18 high-confidence non-indigenous species out of 140, are shared between the Northeast Atlantic Ocean, Macaronesia and the Mediterranean Sea (Fig. 6A). The broad distribution of these non-indigenous species is noteworthy for it reflects a very wide amplitude in abiotic and biotic parameters. At least nine of these widely distributed non-indigenous species (Antithamnion hubbsii/nipponicum, Antithamnionella spirographidis, Asparagopsis armata, Bonnemaisonia hamifera, Codium fragile subsp. fragile, Colpomenia peregrina, Dictyota cyanoloma, Grateloupia turuturu, Scytosiphon dotyi) are reported from all five regions. The remaining species have a more restricted distribution range, being only present in the three central regions Macaronesia, Lusitania and the Western Mediterranean Sea (Phycocalidia suborbiculata, Spongoclonium caribaeum, Symphyocladia marchantioides), or being absent from either of the two peripheral regions, i.e. the Eastern Mediterranean Sea (Antithamnionella ternifolia, Sargassum muticum) or Northern Europe (Asparagopsis taxiformis, Rugulopteryx okamurae, Pachymeniopsis gargiuloi).

The number of high-confidence non-indigenous species currently only found in the Mediterranean Sea (66 species) is striking. Including cryptogenic or data deficient species brings this number to 115. However, there is considerable differentiation of non-indigenous species between the Western and Eastern Mediterranean Sea. Only 34 non-indigenous species are shared between both regions, representing 38% of the total non-indigenous species diversity in the

Mediterranean Sea (Fig. 6C). In addition, the fraction of non-indigenous species unique to the
Eastern Mediterranean Sea (27 species) is considerably larger than for the Western Mediterranean
Sea (8 species) (Fig. 6C). The latter pattern is largely the result of dispersal of warm-adapted
species from the Red Sea and by extension the Indo-Pacific Ocean via the Suez Canal. At present,
only a fraction of these have spread to the Western Mediterranean Sea resulting in a higher diversity
of non-indigenous species in the Eastern Mediterranean Sea.

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The non-indigenous species reported from the Thau Lagoon in France, and the Mar Piccolo and Venice Lagoon in Italy are quite distinct compared to those of surrounding Mediterranean waters. A combination of anthropogenic disturbances and intense aquaculture activities (trade and exchanges), more specifically import of shellfish, has resulted in a very high diversity of nonindigenous species in these lagoon systems (66 species), comparable to that of the surrounding Mediterranean Sea locations despite their much smaller area. The non-indigenous species in Mediterranean lagoons have, moreover, more affinities with the Atlantic Ocean than with the Mediterranean Sea, likely to be due to exchanges between shellfish production areas (see below). Nearly half of the non-indigenous species encountered in Mediterranean aquaculture lagoons are not (yet) reported from surrounding Mediterranean water, while 36 are shared with the Northeast Atlantic Ocean. Of these 36 non-indigenous species, 13 have only been recorded in aquaculture lagoons within the Mediterranean Sea. These mainly include species with a Northwest Pacific Ocean origin, e.g. Dasysiphonia japonica, Neopyropia yezoensis and Nitophyllum stellatocorticatum. Differences in the abiotic physico-chemical environment between Mediterranean lagoons and surrounding coastlines probably underlie the failure of these species to spread widely in the Mediterranean Sea. Rugulopteryx okamurae, however, presents a striking counterexample of this trend. The species was collected in the Thau Lagoon for the first time in 2002 (Verlaque et al., 2009). Initially R. okamurae appeared to be rather non-invasive, but in 2015 it was reported from the Strait of Gibraltar (Ocaña et al., 2016; El Aamri et al., 2018; García-Gómez et al., 2020), where the species forms dense stands rapidly overgrowing most native seaweed species. More recently the same alarming invasive behaviour of R. okamurae has been noted in the Marseille area (Ruitton et al., 2021) as well as southwest Portugal (Liulea et al., 2023) and Macaronesia (Faria et al., 2022).

A Northwest Pacific origin of the largest part of the non-indigenous species present in the Mediterranean lagoons and Northern Europe is well established (Fig. 7, Table 1) (Boudouresque *et al.*, 2010). Regular monitoring and surveys demonstrated that many of those species had been first

accidentally introduced in the Mediterranean lagoons (notably the Thau lagoon) and were then transported to the Northeast Atlantic Ocean hitchhiking with oyster transfers (Mineur et al., 2007a). *Undaria pinnatifida* presents a notable exception to this pattern. Following its accidental introduction in the Thau lagoon, the species was deliberately introduced in Brittany for aquaculture purposes (Floc'h et al., 1991) from which it rapidly spread and established itself as one of the dominant non-indigenous species in artificial as well as natural habitats in the Northeast Atlantic Ocean (Voisin et al., 2005; Guzinski et al., 2018). Note that commercial transfers of oysters between the Mediterranean Sea and the Atlantic coasts of France is still fully allowed under French and European regulations, which results in a quasi-continuous series of secondary introduction events. Aquaculture- and fisheries-associated transport (e.g. nets, packing material) between the Atlantic Ocean and Mediterranean Sea is also undoubtedly responsible for the introduction of a range of native Atlantic Ocean species to the Mediterranean lagoons (e.g. Ascophyllum nodosum, Chorda filum, Grateloupia minima, Fucus spiralis and Vertebrata fucoides).

The Mediterranean Sea, Macaronesia and Lusitania are also home to 35 high-confidence non-indigenous species with presumed or established Australasian origin (Fig. 7). Species with Australasian origin are much less represented in Northern Europe, reflecting the warm temperate nature of these species. The introduction vectors for this category of species remain, however, largely elusive. For *Acrothamnion preissii* and *Womersleyella setacea* ship traffic has been suggested as vector based on their first observation close to a major harbour (Livorno, Italy), but accidental release from scientific laboratories and public or private aquaria is also a possibility. As with the notorious case of *Caulerpa taxifolia* (Verlaque *et al.*, 2015), aquarium releases have likely resulted in the introduction of other seaweeds, mainly in the Mediterranean Sea (Vranken *et al.*, 2018).

Despite the abovementioned clear categories of non-indigenous species and associated pathways, for circa one third of the species there is considerable uncertainty regarding the area of origin and the potential dispersal vectors. Complicating identification of native range and vectors even further, population-level molecular studies on several non-indigenous species have unveiled multiple independent introductions possibly involving different vectors (McIvor *et al.*, 2001; Voisin *et al.*, 2005; Geoffroy *et al.*, 2016; Le Cam *et al.*, 2020).

Introduction hotspots

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600 Of the 140 high-confidence non-indigenous species, 65% have been reported for the first time in the 601 Mediterranean Sea, 26% in the Northeast Atlantic Ocean and 9% in Macaronesia. The distributions 602 of the first record of each species in the study area underscore the importance of aquaculture for introductions of seaweeds. The Thau lagoon, with 30 reports of first introductions in the study area, 603 604 is one of the major introduction hotspots. In total 58 species, constituting 32% of the total seaweed 605 diversity or 48-99% of the biomass, have become established in this coastal lagoon (Boudouresque et al., 2010). The Thau lagoon is an important centre of oyster cultivation in the Mediterranean Sea. 606 607 However, the oyster farmers rely on the import of oyster spat produced in other regions because the 608 lagoon is not particularly suitable for oyster reproduction. Since 1977, only spat of Pacific oysters 609 spat produced in the Atlantic Ocean is allowed to be laid in the French Mediterranean lagoons 610 (Verlaque et al., 2007). However, it is likely that some non-official imports from outside of Europe occur, as reported by Verlaque (1996). These continuous transfers across different biogeographic 611 612 regions result in astonishingly high numbers of non-indigenous species. A low native diversity due 613 to the low occurrence of natural hard substrata in lagoons and relatively recent construction of hard substrata for aquaculture purposes, concomitant with transfers of livestock which seed the new 614 substrata, makes these lagoons hotspots for non-indigenous species establishment (Mineur et al., 615 2015). 616 617 The Southeast Mediterranean Sea constitutes another introduction hotspot, which accounts 618 for 24 first reports and a total of 32 non-indigenous species. The inauguration of the Suez Canal in 619 1869 resulted in an open connection between the Northern Red Sea and the Eastern Mediterranean Sea. As a result, more than 500 marine species are believed to have invaded the Mediterranean Sea 620 through the Suez Canal, so-called Lessepsian migrants (Zenetos et al., 2010; Zenetos et al., 2012; 621 Galil et al., 2021). With respect to non-indigenous seaweeds, many species were first reported in a 622 623 series of papers by the Egyptian phycologist Anwar Aleem (Aleem, 1948; Aleem, 1950; Aleem, 624 1951; Aleem, 1993). Recent efforts by Greek, Lebanese, Israeli and Turkish phycologists have 625 expanded the list of Lessepsian seaweeds considerably (e.g. Tsiamis, 2012; Hoffman, 2013; Bitar et al., 2017; Israel & Einay, 2017; Cinar et al., 2021; Galil et al., 2021). Nevertheless, a paucity of 626 627 historic baseline data makes it often difficult to establish the Lessepsian origin of many species or to 628 point to the exact date of introduction. As outlined above, records of species with clear Indo-Pacific 629 affinities which predate the opening of the Suez Canal (e.g. Ganonema farinosum and Acanthophora nayadiformis) still puzzle phycologists. In addition, the identities of many species 630 631 reported for the first time by Aleem and others (e.g. Gracilaria arcuata, G. disticha, Hypnea

flagelliformis, Solieria dura, Spatoglossum variabile) have never been confirmed using molecular markers and are highly uncertain. In general, a detailed understanding of past and contemporary temporal dynamics of seaweed introductions in the Eastern Mediterranean Sea remains a challenge. More than in any other region it remains difficult to link the observation of a new seaweed species with the introduction date. This uncertainty has bearing on the monitoring of migration through the Suez Canal which has been regarded as an ongoing process (Boudouresque, 1999; Por, 2012). The current construction of the new Suez Canal, doubling the capacity of the current corridor, is expected to further increase the influx of Red Sea species (Galil *et al.*, 2015) and contribute to further tropicalisation of the Mediterranean Sea (Bianchi & Morri, 2003; Coll *et al.*, 2010).

Compared to the two Mediterranean Sea introduction hotspots, first records appear less localised in the Northeast Atlantic Ocean. The English Channel (Brittany, southern English coast) and the Scheldt estuary (the Netherlands) are most prominent as introduction hotspots. To what extent this spatial pattern reflects the true locations of primary introductions or whether the locations of first records are biased by the distribution of preferred study areas of phycologists and research institutes is difficult to assess. There is, indeed, a strong correlation between the introduction hotspots (i.e. locations from which a high number of first records for the study area were reported) and the density map of all records of non-indigenous species, which is indicative of high monitoring activities in areas where many non-indigenous species are found. In addition, it is noteworthy that the English Channel and Scheldt estuary are important areas for oyster farming, and many non-indigenous species have been accidentally introduced with oyster transfer from the Mediterranean Sea.

Introduction rates: temporal trends

Disentangling the factors underpinning temporal trends in the accumulation rate of non-indigenous seaweeds may improve our understanding of introductions and result in better-informed predictions of future trajectories (Seebens *et al.*, 2018). Deducing temporal trends in the rate of introduction of non-indigenous species, however, assumes a correlation between the date of introduction and the moment the species was detected. Although seemingly straightforward, for several species the timespan between introduction and detection is probably considerable and unpredictable. Detection obviously depends on collecting effort, but as highlighted in previous sections, the taxonomic and biogeographic framework will also determine if a species is considered non-indigenous.

Furthermore, Costello & Solow (2003) demonstrated that an increasing rate of detection need not imply an increasing rate of introductions even when collection effort is constant. Given this complexity, reports that introduction rates have increased or decreased in specific time windows should be treated with caution.

Acknowledging this uncertainty, the detection of non-indigenous species shows two distinct phases, one prior to 1950-1970 characterised by low accumulation rates, followed by another much higher accumulation rate from then onward (Fig. 8). Irrespective of taxonomic and biogeographic uncertainties, there is little to no indication for a decline in the rate at which non-indigenous species are reported. The observation that the detection, and presumably also the introduction, of non-indigenous species has not reached saturation (Fig. 8), is in line with the observations by Seebens *et al.* (2017) for other taxonomic groups. For seaweeds, the absence of a decline in the first-record rate may point to the inefficiency of measures aimed to prevent and mitigate new introductions. Alternatively, the community of phycologists involved in monitoring of non-indigenous seaweeds has become larger and more efficient at detecting incoming species, e.g. through the use of DNA-barcoding methods.

According to Mineur et al. (2007b) who sampled the hulls of several commercial cargo vessels, hull fouling seems to play a relatively minor role in the displacement of seaweeds across the globe. Similarly, ballast water, one of the prime sources of introductions for marine invertebrates and microalgae (Bolch & de Salas, 2007; Gollasch et al., 2015), is relatively unimportant with respect to introduction of seaweeds. Yet there is evidence of introduction with commercial ships not hulls or ballast water but other components such as anchors, for instance in Australia for *U. pinnatifida* (South et al., 2017). Leisure boats, however, likely contribute to the local spread of non-indigenous species within the study area. Their role has certainly been underestimated so far, at least for secondary introductions (Mineur et al., 2008). Contrary to the Eastern Mediterranean Sea where the opening of the Suez Canal resulted in an ongoing and steady influx of non-indigenous species (Galil et al., 2015), in the Western Mediterranean Sea a disproportionate number of non-indigenous seaweed species appears to have been introduced through import of oyster stocks (Verlaque et al., 2007). In the late 1960s and early 1970s, disease caused by Asian oysters importation outbreaks in the study area affecting oyster populations caused a major disruption of production (Mineur et al., 2015). Mitigation procedures involved massive imports in the 1970s of Pacific oyster from its native range in the Northwest Pacific Ocean, or via from the Puget Sound in the Northeast Pacific Ocean where the species is also cultivated on a large scale (Mineur et al., 2014). Sargassum muticum introduced into the study area was shown to have

several origins including a primary introduction from Asia and a secondary introduction from the Northeast Pacific Ocean (Le Cam *et al.*, 2020).

Conclusion and perspectives

In conclusion, our critical synthesis of non-indigenous seaweed diversity in the Northeast Atlantic Ocean, Mediterranean Sea and Macaronesia revealed widespread taxonomic and biogeographic uncertainty. This finding negatively impacts efforts to evaluate the effectiveness of measures to reduce non-indigenous species influx, manage their risks and impacts, and devising potential control strategies. This uncertainty can be addressed through the progressive use of molecular markers, particularly standard DNA barcoding approaches, which can in most cases confirm the identification of presumed non-indigenous seaweeds (Viard & Comtet, 2015). Importantly, however, a taxonomic and biogeographic reference framework should also be established for putative indigenous regions. DNA-based identification is relatively well-developed for species with a Northwest Pacific origin. For tropical taxa such a framework lags behind, resulting in higher levels of uncertainty regarding the identity of non-indigenous species with presumed tropical origin. Such a reference framework will also be necessary for early detection of non-indigenous species and monitoring with the use of bulk sample and eDNA metabarcoding (Darling *et al.*, 2017; Keck & Altermatt, 2023).

However, standard DNA barcoding may not be sufficient to interpret more complex introduction histories, such as cases where non-indigenous and indigenous populations co-occur, as suggested for several species in the Mediterranean Sea (Verlaque *et al.*, 2015) or where recent human-mediated dispersal needs to be evaluated against ongoing and natural dispersal events linked to Pleistocene or Holocene climatic oscillations (Neiva *et al.*, 2016). In these cases a combination of population-level sampling strategies and molecular markers that capture intraspecific diversity is needed to shed light on the number and directions of dispersal events and re-evaluate the status of taxa currently considered cryptogenic.

In parallel with DNA-barcoding efforts, historical and reliable baseline data of seaweed diversity is needed to reduce the subjective interpretation of non-indigenous species and determine more precisely their date of introduction. It is likely that the introduction of many non-indigenous species occurred significantly earlier than the time of their first detection, especially for species that have morphologically similar congeners in the study area (such as species from the genera *Dictyota*,

727 Gracilaria, Polysiphonia and Ulva). Herbarium collections can serve as a crucial source of primary data to address this issue, and advances in sequencing technologies make it possible to obtain 728 729 genetic data from voucher specimens tens or even hundreds of years old as demonstrated for the 730 Codium fragile-complex (Provan et al., 2008). We anticipate that herbaria will play an increasingly important role in documenting spatio-temporal patterns of non-indigenous seaweeds, alongside 731 732 large-scale digitization efforts for these collections. More precise estimates of the date of 733 introduction will also reduce uncertainty regarding the accumulation rate of non-indigenous species. 734 Our analyses suggest that the rate of introduction in the study area has not decreased. However, it is 735 unclear whether this trend reflects a steady accumulation of non-indigenous species or increased 736 and more efficient detection through monitoring or advanced identification methods. Based on 737 reliable baselines, temporal surveys will allow to uncover the trends in non-indigenous species introduction rates. Diversifying the type of survey, from punctual surveys (e.g. Bio-Blitzes or Rapid 738 Assessment surveys) to systematic comprehensive surveys (e.g. full inventories), including 739 740 morphological-based and DNA-based assessments, such as metabarcoding (see above), is a need for effective prevention and early detection and also to monitor trends over time in new species 741 742 introductions.

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Our study also reveals significant differences in the geographical distribution of nonindigenous seaweed species across the study area, with only 18 species shared between the three main regions. Non-indigenous species distribution reflects both their abiotic niche and the efficiency of primary and secondary dispersal vectors. It is expected that current patterns will become increasingly homogenous with time due to various factors. First, recently introduced species may continue to expand as part of the ongoing invasion process. Secondly, evolutionary processes such as selection, genetic admixture and hybridisation may occur during the invasion process, leading to adaptation and expansion of the species in new ecological conditions. Last, further range expansions are expected for non-indigenous species with affinities for warm temperate to tropical temperatures as a result of ongoing ocean warming. This could result in an influx of warm-water adapted species into regions where conditions are currently unfavourable, while warming may also render regions unfavourable for non-indigenous species that currently thrive there. It is anticipated that under ocean warming, Eastern Mediterranean Sea non-indigenous species would likely expand their range in western direction, while species ranges in Lusitania would shift northward. The accurate estimation of the rate at which non-indigenous species expand their ranges, either as invasion fronts or more erratically by jump dispersals, rely on detailed

monitoring. Such estimates can inform on dispersal vectors at various spatial scales and guide policy makers to take effective measures to prevent or limit the spread of these species.

We advocate that combining the efforts of taxonomists who provide a reliable framework on the number and likely geographic origin of non-indigenous seaweeds, together with environmental monitoring, offers the best strategy to identify species of concern, characterise their life history traits, and develop effective management strategies. Especially for the species entering the Mediterranean Sea through the Suez Canal, a combination of horizon scanning exercises, intensive monitoring and rapid-response eradication efforts at the local level may be the only tools to try and control the establishment of Lessepsian migrants. With respect to non-indigenous species that hitchhike with shellfish transport, effectively limiting imports into Europe and controlling translocations between regions in Europe should effectively reduce the rate of primary and secondary introductions (Mineur et al., 2014). Alternatively, immersion for shorter periods (3 seconds) at temperatures of 80–85°C is effective in killing macroalgal propagules (Mineur et al., 2007a). In addition, hull fouling has an important role for introductions, especially towards secondary spread of already introduced non-indigenous seaweeds (Clarke Murray et al., 2011). The related guidance developed in the context of the Marine Environmental Protection Committee (MEPC, 2011) is a step forward, but we stress the need for more enforceable control of this pathway. Mitigating the negative effects of non-indigenous species that have already established will likely prove to be even more difficult. Solid baseline data will allow us to detect introduction patterns and non-indigenous species range shifts in early stages and act accordingly.

Combatting the effects of non-indigenous seaweeds will require coordinated action at the European and international level to prevent the introduction of species, to quickly detect and rapidly eradicate species to prevent them from establishing, and to manage established species to minimise their ecological and economic impact (IAS Regulation (EU) 1143/2014 on invasive alien species). These measures will require significant efforts and collaboration between science, management, policy, and society. Our dataset supports these regulatory actions by providing a solid baseline on non-indigenous seaweeds. This baseline contributes to the assessment of the current situation, helps authorities to identify new introductions and monitor the status of already established species, and importantly identifies the current knowledge gaps concerning taxonomic and biogeographic uncertainties.

Acknowledgments

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Disclosure statement

No potential conflict of interest was reported by the authors.

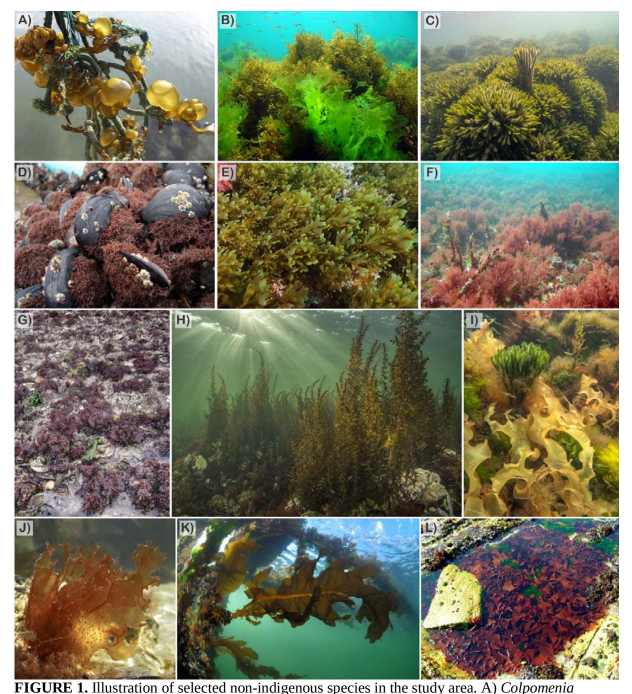
806 Supplementary information

- 807 Supplementary Table S1: concise description of the status of species flagged as non-indigenous in
- 808 the Northeast Atlantic Ocean, the Mediterranean Sea, and Macaronesia.
- 809 The complete dataset containing all records is available at Zenodo (DOI:10.5281/zenodo.7798640).

Author contributions

- 812 LM. van der Loos + Q. Bafort +S. Bosch: concept, data acquisition, analyses, writing; F. Leliaert +
- 813 O. De Clerck: concept, data acquisition, analyses, writing. Other authors: data acquisition, writing.

- 815 **Table 1.** Overview of the non-indigenous seaweed species (Charophyta, Chlorophyta,
- Phaeophyceae, Xanthophyceae, Rhodophyta) reported from the Northeast Atlantic Ocean, the
- 817 Mediterranean Sea, and Macaronesia.



peregrina growing attached to nylon fishing net in a harbour (photo: Frank Perk, the Netherlands);
B) The green non-indigenous species *Ulva australis* (photo: Ignacio Bárbara, Atlantic coast Spain);
C) A dense *Codium fragile* subsp. *fragile* reef (photo: Mick Otten, the Netherlands); D)
Caulacanthus okamurae often grows high in the intertidal (photo: Ignacio Bárbara, Atlantic coast Spain); E) Rugulopteryx okamurae has been introduced in the Northeast Atlantic Ocean,
Mediterranean Sea and in Macaronesia (photo: Sandrine Ruitton, Mediterranean coast France); F)
Asparagopsis armata is often regarded as a high-nuisance invasive species (photo: Ignacio Bárbara,

Atlantic coast Spain); G) Gelidium vagum has only been reported from the Netherlands but is
locally very abundant (photo: Mart Karremans, the Netherlands); H) A Sargassum muticum forest
(photo: Rob Aarsen, the Netherlands); I) Grateloupia turuturu, Codium fragile subsp. fragile, and
Sargassum muticum covering the seabed (photo: Ad Aleman, the Netherlands); J) A fertile
specimen of Nitophyllum stellatocorticatum (photo: Mart Karremans, the Netherlands); K) Undaria
pinnatifida growing attached to aquaculture facilities (photo: Ron Offermans, the Netherlands); L)
A tidal pool with Grateloupia turuturu (photo: Ignacio Bárbara, Atlantic coast of Spain).

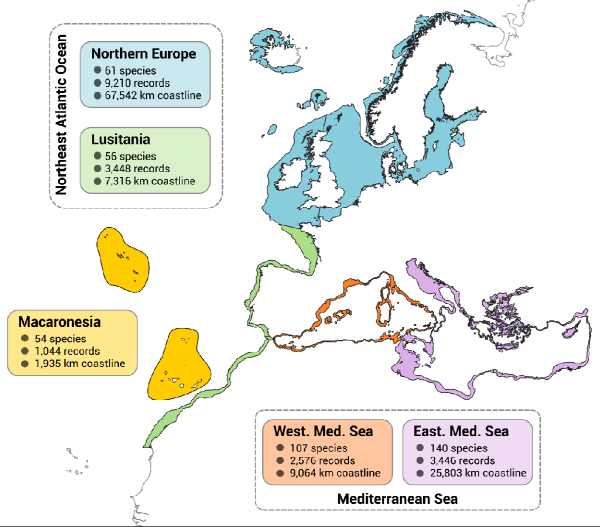


FIGURE 2. Flowchart for assessing the taxonomic confidence and biogeographic status of putative non-indigenous species. This figure builds on the concepts from Essl *et al.* (2018).

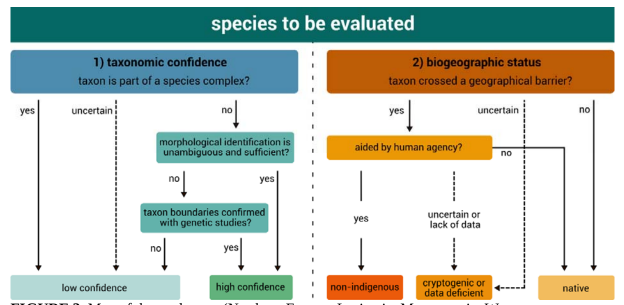


FIGURE 3. Map of the study area (Northern Europe, Lusitania, Macaronesia, Western Mediterranean Sea, and Eastern Mediterranean Sea), with indication of the number of recorded non-indigenous species, number of records and the length of the coastline. The number of species includes cryptogenic and data deficient species.

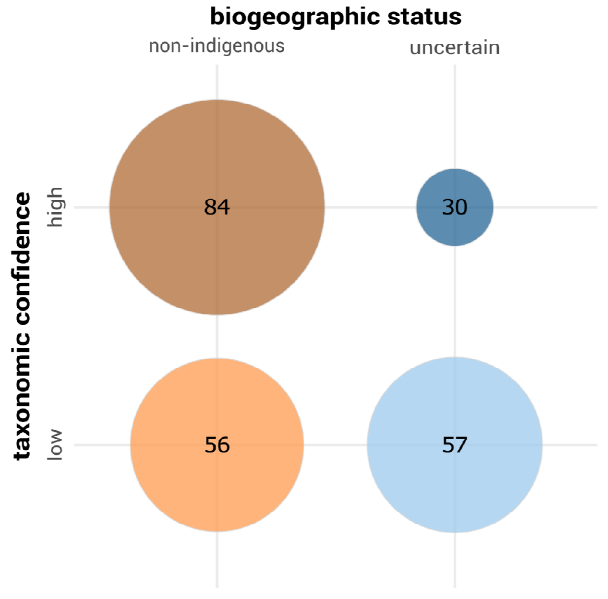


FIGURE 4. Biogeographic status (either non-indigenous or uncertain, the latter category including the 'cryptogenic' and 'data deficient' species) and taxonomic confidence of seaweed species flagged as non-indigenous in the study area. Circle surface area corresponds to the number of species.

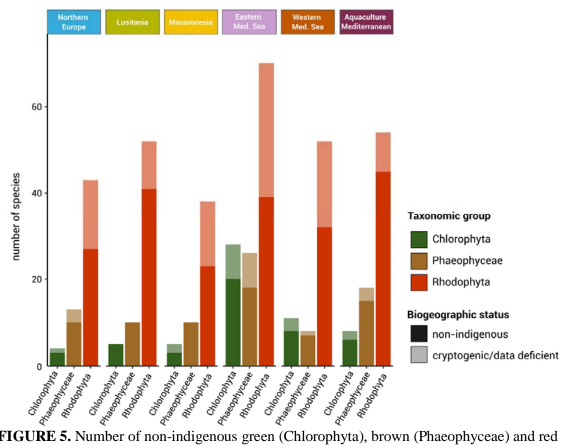


FIGURE 5. Number of non-indigenous green (Chlorophyta), brown (Phaeophyceae) and red (Rhodophyta) non-indigenous species in Northern Europe, Lusitania, Macaronesia, the Eastern and Western Mediterranean Sea, and aquaculture sites in the Mediterranean Sea. Dark shaded colours represent numbers of high-confidence non-indigenous species; light shaded colours represent cryptogenic and data deficient species.

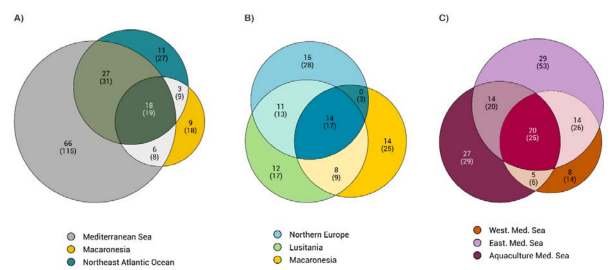
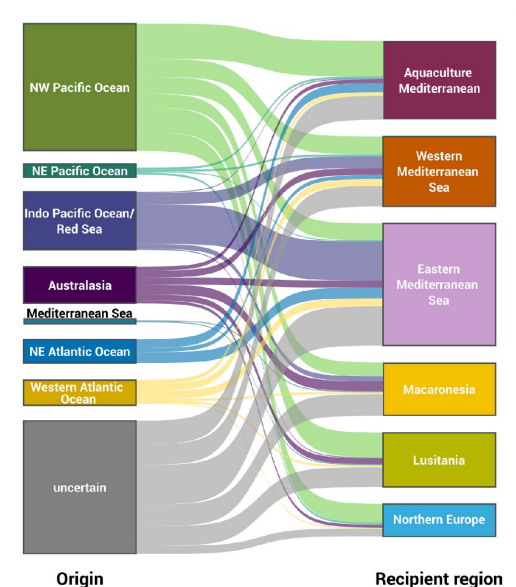


FIGURE 6. Number of non-indigenous species shared among regions. A) Mediterranean Sea, Macaronesia, and Northeast Atlantic Ocean; B) Northern Europe, Lusitania, and Macaronesia; C) Western Mediterranean Sea, Eastern Mediterranean Sea, and aquaculture sites in the Mediterranean Sea. Numbers display the high-confidence non-indigenous species (excluding cryptogenic and data deficient species), with total number of non-indigenous species including cryptogenic and data deficient species in brackets.



860 FIGURE 7.

 The origin of non-indigenous species. For each of the recipient study regions, the Sankey diagram displays what proportion of non-indigenous species have likely originated from a specific area (Northwest Pacific Ocean, Northeast Pacific Ocean, Indo Pacific Ocean/Red Sea, Australasia, Mediterranean Sea, Northeast Atlantic Ocean, Western Atlantic Ocean, or uncertain). The width of the flow arrows is proportional to the number of non-indigenous species (including non-indigenous, cryptogenic and data deficient species).

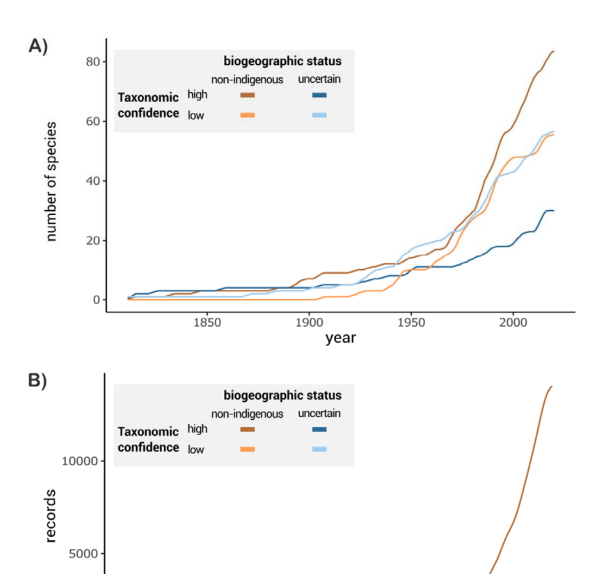


FIGURE 8. The accumulation of A) the number of records, and B) the number of non-indigenous species through time (1808-2022). Colours indicate taxonomic confidence and biogeographic status: dark brown (taxonomic confidence = high, biogeographic status = non-indigenous), light brown (taxonomic confidence = low, biogeographic status = non-indigenous), dark blue (taxonomic confidence = high, biogeographic status = uncertain), light blue (taxonomic confidence = low, biogeographic status = uncertain). All trends are displayed as moving averages over 5 years.

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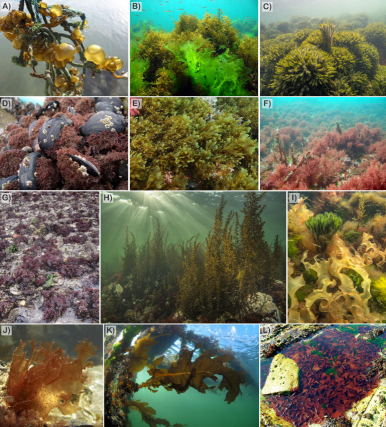
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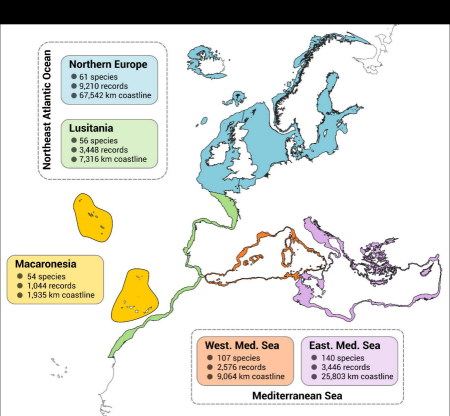
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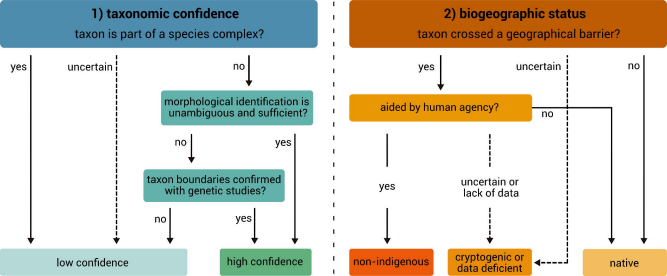
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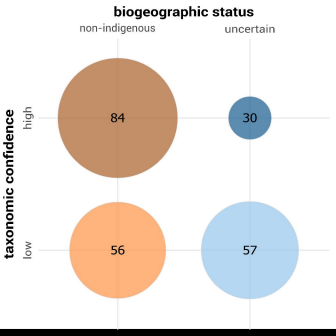
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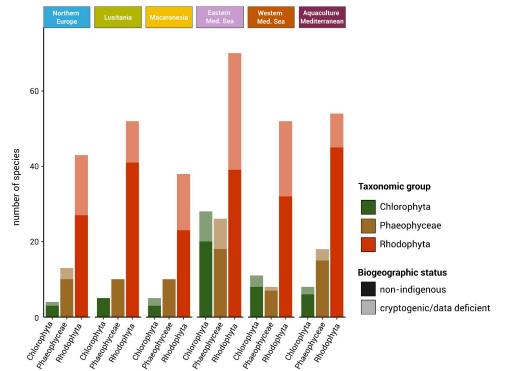


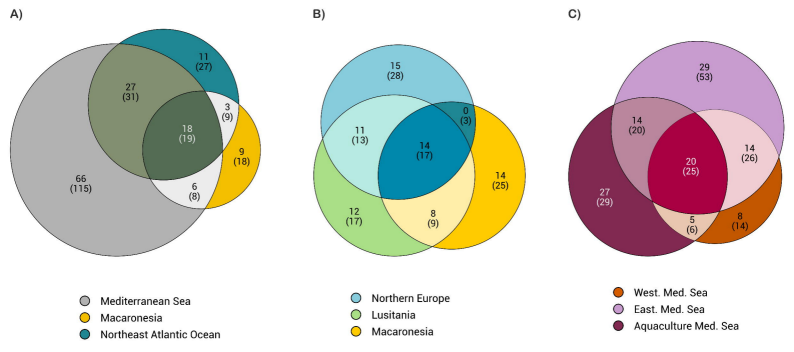


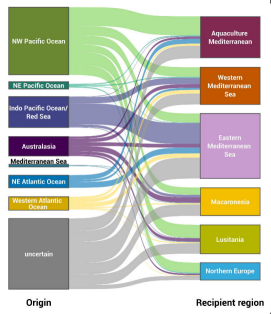
species to be evaluated











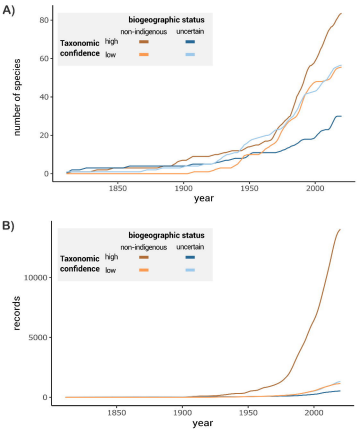


Table 1. Overview of the non-indigenous seaweed species (Charophyta, Chlorophyta, Phaeophyceae, Xanthophyceae, Rhodophyta) reported from the study area.

raiodophyta/ reported fro		date of first recor	d			
a .	Northeast	Mediterranean		Taxonomic	Biogeographical	
Species Charophyta	Atlantic	Sea	Macaronesia	confidence	status	Origin
Chara connivens Salzmann ex A.Braun	native (Baltic Sea: 1870)	native	1975	0	data deficient	uncertain
Chlorophyta	,					
Acetabularia calyculus J.V.Lamouroux	absent	1968	absent	0	cryptogenic	uncertain
Avrainvillea amadelpha (Montagne) A.Gepp & E.S.Gepp	absent	2015	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Batophora occidentalis var. largoensis (J.S.Prince & S.Baker) S.Berger & Kaever ex M.J.Wynne	absent	2020	absent	0	non-indigenous	Western Atlantic Ocean
Bryopsis pennata J.V.Lamouroux	absent	1961	native	0	data deficient	uncertain
Caulerpa chemnitzia (Esper) J.V.Lamouroux	absent	1926	native	0	cryptogenic	Indo Pacific Ocean/Red Sea
Caulerpa cylindracea Sonder	absent	1985	1970	1	non-indigenous	Australasia
Caulerpa denticulata Decaisne	absent	1929	native	1	non-indigenous	Indo Pacific Ocean/Red Sea
Caulerpa integerrima (Zanardini) M.J.Wynne, Verbruggen & D.L.Angel	absent	2020	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
Caulerpa lamourouxii (Turner) C.Agardh	absent	1951	absent	1	cryptogenic	Indo Pacific Ocean/Red Sea
Caulerpa mexicana Sonder ex Kützing	absent	1939	native	0	non-indigenous	Indo Pacific Ocean/Red Sea
Caulerpa prolifera (Forsskål) J.V.Lamouroux	absent	native	native (Azores: 2013)	1	cryptogenic	uncertain
Caulerpa taxifolia (M.Vahl) C.Agardh	absent	1984	native	1	non-indigenous	Australasia
Caulerpa taxifolia var. distichophylla (Sonder) Verlaque, Huisman & Procaccini	absent	2003	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
Caulerpa webbiana Montagne	absent	absent	native (Azores: 2002)	1	cryptogenic	uncertain
Cladophora patentiramea (Montagne) Kützing	absent	1991	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Cladophoropsis fasciculata (Kjellman) Wille	absent	1928	absent	0	data deficient	uncertain

Codium arabicum Kützing	2003	2007	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
Codium fragile subsp. fragile	1845	1946	1990	1	non-indigenous	Northwest Pacific Ocean
Codium parvulum (Bory ex Audouin) P.C.Silva	absent	2004	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
Codium pulvinatum M.J.Wynne & R.Hoffman	absent	2014	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
<i>Codium taylorii</i> P.C.Silva	2004	1939	native	0	non-indigenous	uncertain
Derbesia boergesenii (M.O.P.Iyengar & Ramanathan) Mayhoub	absent	1972	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
<i>Derbesia rhizophora</i> Yamada	absent	1984	absent	0	non-indigenous	Northwest Pacific Ocean
Flabellia petiolata (Turra) Nizamuddin	2013	native	native	1	cryptogenic	uncertain
Halimeda incrassata (J.Ellis) J.V.Lamouroux	absent	2011	2005	1	non-indigenous	Western Atlantic Ocean
Lychaete herpestica (Montagne) M.J.Wynne	absent	1944	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Neomeris annulata Dickie	absent	2003	absent	1	cryptogenic	Indo Pacific Ocean/Red Sea
Parvocaulis parvulus (Solms-Laubach) S.Berger, Fettweiss, Gleissberg, Liddle, U.Richter, Sawitzky & Zuccarello	absent	1930	native	1	cryptogenic	uncertain
Pseudocodium okinawense E.J.Faye, M.Uchimura & S.Smimada	absent	2017	absent	1	non-indigenous	uncertain
Siphonocladus tropicus (P.Crouan & H.Crouan) J.Agardh	absent	2014	native	1	non-indigenous	Western Atlantic Ocean
Ulva australis Areschoug	1990	1984	absent	1	non-indigenous	uncertain
Ulva californica Wille	1999	2011	absent	0	non-indigenous	Northeast Pacific Ocean
<i>Ulva chaugulii</i> M.G.Kavale & M.A.Kazi	absent	2015	absent	1	cryptogenic	Indo Pacific Ocean/Red Sea
Ulva lactuca Linnaeus	absent	1813	absent	1	cryptogenic	uncertain
<i>Ulva ohnoi</i> M.Hiraoka & S.Shimada	absent	2002	absent	1	non-indigenous	Northwest Pacific Ocean
<i>Ulva tepida</i> Y.Masakiyo & S.Shimada	absent	2002	absent	1	non-indigenous	Northwest Pacific Ocean
Ulvaria obscura (Kützing) Gayral ex Bliding	native	1985	absent	0	non-indigenous	uncertain
Uronema marinum Womersley	absent	2008	absent	0	cryptogenic	uncertain

Phaeophyceae						
Acrothrix gracilis Kylin	native	1998	absent	0	non-indigenous	uncertain
Ascophyllum nodosum (Linnaeus) Le Jolis	native	2009	native	1	non-indigenous	Northeast Atlantic Ocean
Botrytella parva (Takamatsu) H.S.Kim	1993	1996	absent	0	non-indigenous	uncertain
Chorda filum (Linnaeus) Stackhouse	native	1981	native	1	non-indigenous	Northeast Atlantic Ocean
Cladosiphon zosterae (J.Agardh) Kylin	native	1985	native	1	non-indigenous	Northeast Atlantic Ocean
Colpomenia peregrina Sauvageau	1905	1918	1965	1	non-indigenous	Northwest Pacific Ocean
Corynophlaea crispa (Harvey) Kuckuck	native	2003	native	0	data deficient	Northeast Atlantic Ocean
Corynophlaea cystophorae J.Agardh	absent	absent	1993	0	non-indigenous	Australasia
Corynophlaea umbellata (C.Agardh) Kützing	1986	native	uncertain	0	cryptogenic	Northwest Pacific Ocean
Corynophlaea verruculiformis (Y P.Lee & I.K.Lee) Y P.Lee	1994	absent	absent	0	non-indigenous	Northwest Pacific Ocean
Cutleria multifida (Turner) Greville	native	1950	native	1	cryptogenic	Northwest Pacific Ocean
Desmarestia viridis (O.F.Müller) J.V.Lamouroux	native	1947	absent	1	non-indigenous	uncertain
Desmotrichum tenuissimum (C.Agardh) Athanasiadis	native	1947	absent	0	non-indigenous	Northwest Pacific Ocean
Dictyota acutiloba J.Agardh	absent	2010	absent	1	non-indigenous	uncertain
Dictyota cyanoloma Tronholm, De Clerck, A.Gómez-Garreta & Rull Lluch	1995	1935	2006	1	non-indigenous	Australasia
Ectocarpus siliculosus var. hiemalis (P.Crouan & H.Crouan ex Kjellman) Gallardo	native	1985	absent	0	cryptogenic	Northeast Atlantic Ocean
Fucus distichus subsp. evanescens (C.Agardh) H.T.Powell	native (Oslofjord: 1883)	absent	absent	1	non-indigenous	Western Atlantic Ocean
Fucus serratus Linnaeus	native (Iceland: 1897)	absent	native	1	non-indigenous	uncertain
Fucus spiralis Linnaeus	native	1987	native	1	non-indigenous	Northeast Atlantic Ocean
Halothrix lumbricalis (Kützing) Reinke	native	1978	absent	0	non-indigenous	uncertain

Hydroclathrus tilesii (Endlicher) Santiañez & M.J.Wynne	absent	absent	2016	1	non-indigenous	Northwest Pacific Ocean
Leathesia marina (Lyngbye) Decaisne	native	1905	native	1	cryptogenic	Northwest Pacific Ocean
Lobophora lessepsiana C.W.Vieira	absent	2017	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea
Lobophora schneideri C.W.Vieira	absent	2016	native	1	non-indigenous	Western Atlantic Ocean
Microspongium globosum Reinke	native	2003	absent	0	data deficient	Northeast Atlantic Ocean
Myrionema grateloupiae Noda	2006	absent	absent	0	data deficient	Northwest Pacific Ocean
Padina boergesenii Allender & Kraft	absent	1962	native	0	non-indigenous	Indo Pacific Ocean/Red Sea
Padina boryana Thivy	absent	1974	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Padina tetrastromatica Hauck	absent	2004	absent	0	data deficient	Indo Pacific Ocean/Red Sea
Papenfussiella kuromo (Yendo) Inagaki	absent	absent	1990	0	non-indigenous	Northwest Pacific Ocean
Petalonia binghamiae (J.Agardh)	absent	absent	1980	0	non-indigenous	uncertain
K.L.Vinogradova Pylaiella littoralis (Linnaeus) Kjellman	native	1924	absent	1	cryptogenic	Northwest Pacific Ocean
Rugulopteryx okamurae (E.Y.Dawson) I.K.Hwang, W.J.Lee & H.S.Kim	2017	2002	2019	1	non-indigenous	Northwest Pacific Ocean
Saccharina japonica (Areschoug) C.E.Lane, C.Mayes, Druehl & G.W.Saunders	1979	1976	absent	1	non-indigenous	Northwest Pacific Ocean
Sargassum latifolium (Turner) C.Agardh	absent	1986	absent	0	data deficient	Indo Pacific Ocean/Red Sea
Sargassum muticum (Yendo) Fensholt	1960	1980	2020	1	non-indigenous	Northwest Pacific Ocean
Scytosiphon dotyi M.J.Wynne	1987	1960	1990	0	non-indigenous	Northwest Pacific Ocean
Spatoglossum variabile Figari & De Notaris	absent	1944	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Sphaerotrichia firma (E.S.Gepp) A.D.Zinova	absent	1970	absent	1	non-indigenous	Northwest Pacific Ocean
Stypopodium schimperi (Kützing) Verlaque & Boudouresque	absent	1973	1997	1	non-indigenous	Indo Pacific Ocean/Red Sea
Ulonema rhizophorum Foslie	native	2012	absent	0	data deficient	Northeast Atlantic Ocean

Undaria pinnatifida (Harvey) Suringar	1982	1971	absent	1	non-indigenous	Northwest Pacific Ocean
Xanthophyceae						
Vaucheria longicaulis Hoppaugh	1993	absent	absent	0	cryptogenic	uncertain
Rhodophyta						
Acanthophora muscoides (Linnaeus) Bory	absent	1977	absent	0	data deficient	uncertain
Acanthophora nayadiformis (Delile) Papenfuss	absent	1813	absent	1	cryptogenic	Indo Pacific Ocean/Red Sea
Acanthosiphonia echinata (Harvey) Savos &	absent	2018	absent	1	non-indigenous	Western Atlantic Ocean
G.W.Saunders Acrochaetium balticum (Rosenvinge) Aleem & Schulz	1998	absent	absent	0	cryptogenic	Northeast Atlantic Ocean
Acrochaetium catenulatum M.Howe	1967	absent	absent	0	cryptogenic	uncertain
Acrochaetium spathoglossi Børgesen	absent	1944	absent	0	cryptogenic	uncertain
Acrochaetium subseriatum Børgesen	absent	1944	absent	0	cryptogenic	uncertain
Acrothamnion preissii (Sonder) E.M.Wollaston	absent	1969	2009	1	non-indigenous	Indo Pacific Ocean/Red Sea
Agardhiella subulata (C.Agardh) Kraft & M.J.Wynne	1973	1984	absent	0	non-indigenous	uncertain
Aglaothamnion cordatum (Børgesen) Feldmann-Mazoyer	absent	native	2006	1	cryptogenic	uncertain
Aglaothamnion feldmanniae Halos	native	1975	native	0	non-indigenous	uncertain
Aglaothamnion halliae (Collins) Aponte, D.L.Ballantine & J.N.Norris	1960	2017	absent	0	non-indigenous	Western Atlantic Ocean
Ahnfeltiopsis flabelliformis (Harvey) Masuda	absent	1994	absent	0	non-indigenous	Northwest Pacific Ocean
Anotrichium furcellatum (J.Agardh) Baldock	1914	1926	1930	0	cryptogenic	uncertain
Antithamnion amphigeneum A.Millar	1995	1989	absent	1	non-indigenous	Australasia
Antithamnion densum (Suhr) M.Howe	1968	absent	1990	0	non-indigenous	uncertain
Antithamnion diminuatum Wollaston	absent	absent	1988	1	non-indigenous	Australasia
Antithamnion hubbsii/nipponicum	2003	1988	1989	1	non-indigenous	uncertain
Antithamnionella boergesenii (Cormaci & G.Furnari) Athanasiadis	2004	1937	native	0	cryptogenic	Western Atlantic Ocean

Antithamnionella elegans (Berthold) J.H.Price & D.M.John	1961	1882	uncertain	0	cryptogenic	uncertain
Antithamnionella spirographidis (Schiffner)	1931	1905	1974	1	non-indigenous	uncertain
E.M.Wollaston Antithamnionella sublittoralis (Setchell & N.L.Gardner) Athanasiadis	absent	1980	absent	0	cryptogenic	Northeast Pacific Ocean
Antithamnionella ternifolia (Hooker f. & Harvey) Lyle	1906	1926	2005	0	non-indigenous	Australasia
Asparagopsis armata Harvey	1922	1923	1928	1	non-indigenous	Australasia
Asparagopsis taxiformis (Delile) Trevisan	2000	1813	1840	1	non-indigenous	Australasia
Bonnemaisonia hamifera Hariot	1893	1909	1921	1	non-indigenous	Northwest Pacific Ocean
Botryocladia madagascariensis G.Feldmann	absent	1991	1988	0	non-indigenous	Indo Pacific Ocean/Red Sea
Botryocladia wrightii (Harvey) W.E.Schmidt, D.L.Ballantine & Fredericq	2002	1978	absent	1	non-indigenous	Northwest Pacific Ocean
Calliblepharis rammediorum R.Hoffman, M.J.Wynne & G.W.Saunders	absent	2013	absent	1	cryptogenic	uncertain
Caulacanthus okamurae Yamada	1986	2004	absent	1	non-indigenous	Northwest Pacific Ocean
Ceramium atrorubescens Kylin	absent	absent	1988	0	non-indigenous	Indo Pacific Ocean/Red Sea
Ceramium bisporum D.L.Ballantine	absent	1980	absent	0	cryptogenic	Western Atlantic Ocean
Ceramium camouii E.Y.Dawson	absent	2020	absent	0	data deficient	uncertain
Ceramium cingulatum Weber Bosse	absent	absent	1991	0	cryptogenic	uncertain
Ceramium graecum Lazaridou & Boudouresque	absent	1990	absent	0	cryptogenic	uncertain
Ceramium strobiliforme G.W.Lawson & D.M.John	absent	1990	absent	0	non-indigenous	uncertain
Ceramium sungminbooi J.R.Hughey & G.H.Boo	1990	absent	absent	1	non-indigenous	Northwest Pacific Ocean
Champia compressa Harvey	absent	2012	absent	0	data deficient	uncertain
Chondracanthus sp.	2009	absent	absent	1	non-indigenous	uncertain
Chondria coerulescens (J.Agardh) Sauvageau	native	1973	native	1	cryptogenic	Northeast Atlantic Ocean

Chondria curvilineata Collins & Hervey	absent	1980	absent	0	non-indigenous	Western Atlantic Ocean
Chondria polyrhiza Collins & Hervey	absent	1987	absent	0	data deficient	Western Atlantic Ocean
Chondria pygmaea Garbary & Vandermeulen	absent	1974	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Chondrus giganteus Yendo	absent	1994	absent	1	non-indigenous	Northwest Pacific Ocean
Colaconema codicola (Børgesen) Stegenga, J.J.Bolton & R.J.Anderson	1931	1952	native	0	cryptogenic	uncertain
Colaconema dasyae (Collins) Stegenga, I.Mol, Prud'homme & Lokhorst	1951	absent	absent	0	cryptogenic	uncertain
Colaconema robustum (Børgesen) Huisman & Woelkerling	absent	1944	absent	0	cryptogenic	uncertain
Corynomorpha prismatica (J.Agardh) J.Agardh	absent	absent	1990	1	cryptogenic	Indo Pacific Ocean/Red Sea
Cryptonemia hibernica Guiry & L.M.Irvine	1960	absent	absent	1	non-indigenous	uncertain
Dasya baillouviana (S.G.Gmelin) Montagne	1950	native	native	0	cryptogenic	uncertain
Dasya sessilis Yamada	1989	1984	absent	1	non-indigenous	Northwest Pacific Ocean
Dasysiphonia japonica (Yendo) HS.Kim	1984	1998	absent	1	non-indigenous	Northwest Pacific Ocean
Dichotomaria obtusata (J.Ellis & Solander) Lamarck	absent	2014	native	0	non-indigenous	uncertain
Diplothamnion jolyi C.Hoek	uncertain	2012	uncertain	0	data deficient	Western Atlantic Ocean
Dipterosiphonia dendritica (C.Agardh) F.Schmitz	absent	1979	native	0	data deficient	uncertain
Eutrichosiphonia paniculata (Montagne) D.E.Bustamante & T.O.Cho	absent	1967	absent	0	non-indigenous	uncertain
Ezo epiyessoense Adey, Masaki & Akioka	1983	absent	absent	1	data deficient	Northwest Pacific Ocean
Fredericqia deveauniensis Maggs, L.Le Gall, Mineur, Provan & G.W.Saunders	1980	absent	absent	1	cryptogenic	Western Atlantic Ocean
Galaxaura rugosa (J.Ellis & Solander) J.V.Lamouroux	absent	1990	native	1	non-indigenous	Indo Pacific Ocean/Red Sea
Ganonema farinosum (J.V.Lamouroux) K C.Fan & YC.Wang	absent	1808	native	0	cryptogenic	Indo Pacific Ocean/Red Sea

Gayliella fimbriata (Setchell & N.L.Gardner) T.O.Cho	absent	2013	absent	0	non-indigenous	uncertain
& S.M.Boo <i>Gelidium vagum</i> Okamura	2010	absent	absent	1	non-indigenous	Northwest Pacific Ocean
Goniotrichopsis sublittoralis G.M.Smith	1975	1989	absent	0	cryptogenic	Northeast Pacific Ocean
<i>Gracilaria arcuata</i> Zanardini	absent	1931	absent	0	cryptogenic	Indo Pacific Ocean/Red Sea
Gracilaria disticha (J.Agardh) J.Agardh	absent	1924	absent	0	data deficient	Indo Pacific Ocean/Red Sea
Gracilaria vermiculophylla (Ohmi) Papenfuss	1994	2008	absent	1	non-indigenous	Northwest Pacific Ocean
Gracilariopsis chorda (Holmes) Ohmi	2010	absent	absent	1	non-indigenous	Northwest Pacific Ocean
Grallatoria reptans M.Howe	absent	absent	1988	0	cryptogenic	uncertain
Grateloupia asiatica S.Kawaguchi &	2010	1984	absent	1	non-indigenous	Northwest Pacific Ocean
H.W.Wang Grateloupia gibbesii Harvey	absent	1992	absent	1	non-indigenous	Western Atlantic Ocean
Grateloupia imbricata Holmes	2014	absent	2006	1	non-indigenous	Northwest Pacific Ocean
Grateloupia minima P.Crouan & H.Crouan	native	1998	absent	1	non-indigenous	Northeast Atlantic Ocean
Grateloupia patens (Okamura) Kawaguchi & H.W.Wang	absent	1994	absent	1	non-indigenous	Northwest Pacific Ocean
Grateloupia subpectinata Holmes	1947	1990	absent	1	non-indigenous	Northwest Pacific Ocean
<i>Grateloupia turuturu</i> Y.Yamada	1969	1982	1983	1	non-indigenous	Northwest Pacific Ocean
Grateloupia yinggehaiensis H.W.Wang & R.X.Luan	absent	2008	absent	1	non-indigenous	Northwest Pacific Ocean
Griffithsia corallinoides (Linnaeus) Trevisan	native	1964	absent	0	non-indigenous	Northwest Pacific Ocean
Gymnophycus hapsiphorus Huisman & Kraft	absent	absent	1989	0	non-indigenous	uncertain
Herposiphonia parca Setchell	2005	1997	absent	0	non-indigenous	uncertain
Hypnea anastomosans Papenfuss, Lipkin & P.C.Silva	absent	1972	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Hypnea cervicornis J.Agardh	absent	1926	uncertain	0	non-indigenous	uncertain
Hypnea cornuta (Kützing) J.Agardh	absent	1894	absent	1	non-indigenous	Indo Pacific Ocean/Red Sea

Hypnea corona Huisman & Petrocelli	absent	2000	absent	1	non-indigenous	uncertain
Hypnea flagelliformis Greville ex J.Agardh	absent	1956	2007	0	cryptogenic	uncertain
Hypnea valentiae (Turner) Montagne	absent	1996	native	0	non-indigenous	uncertain
Hypoglossum caloglossoides M.J.Wynne & Kraft	absent	2013	absent	0	non-indigenous	Australasia
Hypoglossum heterocystideum (J.Agardh) J.Agardh	absent	absent	2014	0	cryptogenic	Australasia
Kapraunia schneideri (Stuercke & Freshwater) Savoie & G.W.Saunders	2010	1992	absent	1	non-indigenous	Western Atlantic Ocean
Laurencia brongniartii J.Agardh	1989	absent	1994	0	cryptogenic	uncertain
Laurencia caduciramulosa Masuda & S.Kawaguchi	absent	1991	2006	0	non-indigenous	uncertain
Laurencia okamurae Yamada	absent	1984	absent	1	non-indigenous	Northwest Pacific Ocean
Lithophyllum yessoense Foslie	absent	1994	absent	1	non-indigenous	Northwest Pacific Ocean
<i>Lomentaria flaccida</i> Tak.Tanaka	absent	2002	absent	1	data deficient	Northwest Pacific Ocean
Lomentaria hakodatensis Yendo	1984	1978	absent	1	non-indigenous	Northwest Pacific Ocean
Lophocladia lallemandii (Montagne) F.Schmitz	absent	1900	absent	0	cryptogenic	Indo Pacific Ocean/Red Sea
Lophocladia trichoclados (C.Agardh) F.Schmitz	absent	uncertain	1896	0	cryptogenic	Western Atlantic Ocean
Melanothamnus collabens (C.Agardh) Díaz-Tapia & Maggs	1824	absent	absent	1	cryptogenic	Northwest Pacific Ocean
Melanothamnus harveyi/japonicus	1832	1958	1990	1	non-indigenous	Northwest Pacific Ocean
Melanothamnus pseudoforcipatus Díaz- Tapia	2014	absent	2018	1	cryptogenic	uncertain
Monosporus indicus Børgesen	absent	2015	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Nemalion vermiculare Suringar	absent	2005	absent	1	non-indigenous	Northwest Pacific Ocean
Neoizziella divaricata (C.K.Tseng) SM.Lin, SY.Yang & Huisman	absent	absent	1990	0	cryptogenic	Northwest Pacific Ocean
Neopyropia drachii (Feldmann) J.Brodie	1948	absent	absent	1	cryptogenic	uncertain
Neopyropia koreana (M.S.Hwang & I.K.Lee) LE.Yang & J.Brodie	absent	2000	absent	1	cryptogenic	Northwest Pacific Ocean

Neopyropia leucosticta (Thuret) LE.Yang & J.Brodie	1857	absent	1897	1	cryptogenic	uncertain
Neopyropia yezoensis (Ueda) LE.Yang & J.Brodie	1984	1975	absent	1	non-indigenous	Northwest Pacific Ocean
Nitophyllum stellatocorticatum Okamura	2006	1984	absent	1	non-indigenous	Northwest Pacific Ocean
Osmundea oederi (Gunnerus) G.Furnari	native	1987	native	1	cryptogenic	uncertain
Pachymeniopsis gargiuloi S.Y.Kim, Manghisi, Morabito & S.M.Boo	2010	2000	2007	1	non-indigenous	Northwest Pacific Ocean
Pachymeniopsis lanceolata (Okamura) Yamada ex Kawabata	2019	1982	absent	1	non-indigenous	Northwest Pacific Ocean
Palisada maris-rubri (K.W.Nam & Saito) K.W.Nam	absent	1990	absent	0	cryptogenic	Indo Pacific Ocean/Red Sea
Phrix spatulata (E.Y.Dawson) M.J.Wynne, M.Kamiya	absent	1992	absent	1	non-indigenous	uncertain
& J.A.West Phycocalidia suborbiculata (Kjellman) Santiañez & M.J.Wynne	2010	2010	1993	1	non-indigenous	Northwest Pacific Ocean
Pikea californica Harvey	1967	absent	absent	1	non-indigenous	uncertain
Plocamium ovicorne Okamura	2014	absent	absent	0	non-indigenous	Northwest Pacific Ocean
Plocamium secundatum (Kützing) Kützing	absent	1976	absent	0	non-indigenous	uncertain
Polyopes lancifolius (Harvey) Kawaguchi & Wang	2008	absent	absent	1	non-indigenous	Northwest Pacific Ocean
Polysiphonia atlantica Kapraun & J.N.Norris	native	1969	native	0	cryptogenic	Northeast Atlantic Ocean
<i>Polysiphonia delicata</i> Díaz-Tapia	2014	absent	absent	1	cryptogenic	uncertain
Polysiphonia havanensis Montagne	absent	2012	native	0	cryptogenic	Western Atlantic Ocean
Polysiphonia kampsaxii Børgesen	absent	1986	absent	0	data deficient	Indo Pacific Ocean/Red Sea
Polysiphonia morrowii/senticulosa	1993	1996	absent	1	non-indigenous	uncertain
<i>Polysiphonia radiata</i> Díaz-Tapia	2014	absent	absent	1	cryptogenic	uncertain
Predaea huismanii Kraft	absent	absent	1991	0	non-indigenous	uncertain
Rhodophysema georgei Batters	native	1978	absent	0	non-indigenous	Northeast Atlantic Ocean
Rhodymenia erythraea Zanardini	absent	1948	absent	0	data deficient	Indo Pacific Ocean/Red Sea

Sarconema filiforme (Sonder) Kylin	absent	1944	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Sarconema scinaioides Børgesen	absent	1945	absent	0	data deficient	Indo Pacific Ocean/Red Sea
Scageliopsis patens E.M.Wollaston	2004	absent	1989	0	non-indigenous	Australasia
Schizymenia apoda (J.Agardh) J.Agardh	2013	absent	2004	0	cryptogenic	uncertain
Schizymenia dubyi (Chauvin ex Duby) J.Agardh	native	2008	native	0	non-indigenous	Northeast Atlantic Ocean
Schizymenia jonssonii K.Gunnarsson & J.Brodie	1897	absent	absent	1	cryptogenic	uncertain
Scinaia acuta M.J.Wynne	absent	absent	1989	0	non-indigenous	Australasia
Solieria chordalis (C.Agardh) J.Agardh	native (British Isles: 1977)	native	absent	0	cryptogenic	uncertain
Solieria dura (Zanardini) F.Schmitz	absent	1944	absent	0	non-indigenous	Indo Pacific Ocean/Red Sea
Solieria filiformis (Kützing) Gabrielson	2005	1922	native	0	non-indigenous	uncertain
Solieria sp.	2005	2011	absent	1	non-indigenous	Northwest Pacific Ocean
Spermothamnion cymosum (Harvey) De Toni	absent	2008	absent	1	non-indigenous	Australasia
Spongoclonium caribaeum (Børgesen) M.J.Wynne	1973	1974	1980	0	non-indigenous	uncertain
Spyridia aculeata (C.Agardh ex Decaisne) Kützing	native	1937	native	1	cryptogenic	uncertain
Symphyocladia marchantioides (Harvey) Falkenberg	2004	1984	1971	1	non-indigenous	uncertain
Symphyocladiella dendroidea (Montagne) D.Bustamante, B.Y.Won, S.C.Lindstrom & T.O.Cho	2005	1993	absent	1	non-indigenous	uncertain
Vertebrata fucoides (Hudson) Kuntze	native	1988	native	1	cryptogenic	Northeast Atlantic Ocean
Womersleyella setacea (Hollenberg) R.E.Norris	absent	1986	1983	0	cryptogenic	uncertain
Xiphosiphonia pinnulata (Kützing) Savoie & G.W.Saunders	native (British Isles: 1990)	native	2006	0	data deficient	Mediterranean