Species delimitation of eight ascidian (Tunicata) species from the North Eastern Mediterranean Arzu Karahan^a, Berivan Temiz^b, Esra Öztürk^a, Jacob Douek^c, Baruch Rinkevich^c ^aMiddle East Technical University, Institute of Marine Science, Department of Marine Biology and Fisheries, Mersin, Turkiye ^bDevelopmental Biology and Genomics Laboratory, Department of Anatomy, Otago School of Medical Sciences, University of Otago, P.O. Box 56, Dunedin 9054, New Zealand ^cIsrael Oceanographic and Limnology Research, National Institute of Oceanography, Tel Shikmona, PO Box 9753, Haifa 3109701, Israel Corresponding author: Arzu Karahan, Middle East Technical University, Institute of Marine Sciences, Erdemli, Mersin, Turkiye. E-mail: arzukarahan@ims.metu.edu.tr, arzuukarahan@gmail.com, Tel: +90 532 6462547. https://orcid.org/0000-0002-4096-9372

Abstract

Members of the tunicates, a subphylum of marine filter-feeder chordates, inhabit all marine and oceanic habitats from the subtidal to the abyssal. Considered as the closest relatives to the vertebrates, the tunicates are widely used as model organisms for evo-devo, allorecognition, senescence, and whole-body regeneration studies. Yet, species boundaries are poorly understood due to high morphological and genetic plasticity that characterize many tunicates taxa. Here we study taxonomy and the distribution of eight tunicate species (Botrylloides niger Herdman, 1886/ aff. leachii, Botrylloides israeliense Brunetti, 2009, Botrylloides sp., Botrylloides anceps (Herdman, 1891), Botryllus schlosseri (Pallas, 1766), Didemnum perlucidum Monniot F., 1983, Symplegma brakenhielmi (Michaelsen, 1904) and Polyclinum constellatum Savigny, 1816) sampled from six Turkish North Eastern Mediterranean Sea sites and employed the mitochondrial barcoding marker (COI) for evaluating the relationships among geographically restricted and widely spread ascidian species. Species delimitation analysis was conducted using NCBI and the present study sequences. Morphological examinations were first done in the field and then, styelide colonies were cultured in the laboratory and studied using stereo and light microscopes. A putative new *Botrylloides* species (Botrylloides sp.) from the Antalya region was revealed, with 99% matching on the COI gene from Saudi Arabia, further awaiting for detailed traditional taxonomy.

Keywords: Botryllid ascidians, *Didemnum*; *Polyclinum*; *Symplegma*, COI, species delimitation; Turkiye, Mediterranean Sea.

Introduction

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The ascidians (Phylum: Chordata, Subphylum; Tunicata) are a class of marine filter feeder organisms with ca. 3000 described species (https://www.marinespecies.org/ascidiacea) that inhabit all marine and oceanic habitats from the subtidal to the abyssal zone. As the closest relatives of the vertebrates, the tunicates are widely used as model organisms for evo-devo research analyses, for elucidating the evolution of immunity, senescence and ageing processes, for whole-body regeneration phenomena, stem cell biology, and more. Yet, species boundaries for many clades are poorly understood due to high genetic and morphological plasticity, revealing high cases of cryptic diversity (Burnet, 1971; Rinkevich et al., 1993; Denoeud et al., 2010; Reem et al., 2017, 2022; Viard et al., 2019). Further, ascidians' classification requires experienced taxonomists (Rubinstein et al., 2013), a vanishing scientific discipline, in addition to the difficulties in assigning differentiating taxonomic characteristics between closely related species (Rocha et al., 2012). Nowadays, in addition to classical taxonomy, researchers use a wide range of molecular tools to elucidate biodiversity and to solve emerging species delineation issues (Karahan et al., 2016; Brunetti et al., 2017; Viard et al., 2019; Reem et al., 2017, 2022). The cytochrome oxidase subunit 1 (COI) gene is one of the most commonly used markers for this purpose (Hebert et al., 2003). Studies on the taxonomy and distribution of Mediterranean Sea ascidians, now almost bicentennial old (Schlosser and Ellis, 1756; Spallanzani & Chiereghin, 1784; Savigny, 1816) were concentrated primarily on western European coasts and seas, neglecting the eastern basin areas (Berrill, 1950; Rinkevich et al., 1993). One example is the Levant, an area experiencing a continuous flow of non-indigenous species (NIS), including new exotic tropical ascidian species, entering the Mediterranean Sea through the Suez Canal (López-Legentil et al., 2015; Zenetos et al., 2017; Galil et al., 2018). Another less studied area is the Turkish Mediterranean coastline, where the knowledge on ascidians has improved considerably in the past vicenary. So far, about 50 ascidian species (native and NIS), consisting primarily of solitary species, have been recorded from all over the Turkish coastlines (Uysal, 1976; Çınar et al., 2006; Okuş et al., 2007; Çınar, 2014), in comparison to 45 ascidian species reported from the Levantine basin, along the shores of Israel, Egypt, and the Gulf of Suez (Koukouras et al., 1995; Halim & Abdel Messeih, 2016; Reem et al., 2017). For updating some of the less studied ascidian species in the Turkish Mediterranean coasts, this study aimed in clarifying the taxonomy, and further apprising the distributions of eight ascidian species by employing COI analyses and major morphological features.

Materials and Methods

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Sampling and general morphological examinations

Specimens were collected from patchy stony-rocky areas laying on sandy and shallow bottoms (underneath stones, <1 m depth) using razor blades in 6 sites along the Mediterranean coastline of Turkiye, placed, each, in 1,5 mL tube containing 70% ethanol, and kept in room temperature until use (Table S1, Fig. 1). Other colonial fragments were kept in 4% formaldehyde solution at room temperature for morphological analyses. Specimens of all Styelidae species (genera *Botryllus, Botrylloides* and *Symplegma*) were cultured on microscope slides in the Institute of Marine Sciences-Middle East Technical University (IMS-METU) aquaculture room according to Karahan *et al.*, (2022) and used for morphological examination. Morphological examinations were carried out for major morphological features (zooid distributions, colonial structures, colors, spicule shape, oral tentacles, blastogenic life cycles), first by the naked eye, followed by stereo and light microscopes (Olympus SZX16 - UC30 camera; Olympus CX43-ToupTek camera).

DNA extraction, Polymerase Chain Reactions (PCR), and data analysis

Total DNA was extracted from colony fragments according to a modified phenol-chloroform method (Karahan et al., 2022). Isolated DNA was quantified using a Nanodrop spectrophotometer and diluted, when needed, to 20 ng/µl. PCR were performed in 50 µL total volume with 0.5 μM forward and reverse primers and around 10-20 ng/μl of DNA in a readyto-use PCR Master Mix (Thermo Scientific) on the mitochondrial cytochrome oxidase subunit I (COI) using Reem al. (2017)primers (F2gene, et R2-'AMWAATCATAAAGATATTRGWAC'-3 and AARAARGAMGTRTTRAAATTHCGATC'-3). The PCR products were purified and

117 AARAARGAMGTRTTRAAATTHCGATC'-3). The PCR products were purified and

sequenced for forward and reverse directions by Macrogen Inc. (Seoul, South Korea). Detailed

data for voucher specimens' DNA (held in the IMS-METU genetic laboratory) were uploaded

to the Barcode of Life Data System (BOLD, http://www.boldsystems.org; Table S2).

121 BLAST analysis was performed using the GenBank (http://blast.ncbi.nlm.nih.gov/Blast.cgi)

and BOLD engines. Sequences were translated and aligned via MAFFTv7 (Katoh et al., 2018)

and trimmed using Jalview (v 2.11.1.7, Waterhouse *et al.*, 2009). The best model for the MrBayes (Ronquist *et al.*, 2012) was chosen via PhyML-SMS v3 software (SMS: Smart Model Selection in PhyML; Lefort *et al.*, 2017). MrBayes were run according to the GTR+R model for 10.800.000 combined states (two independent runs), resulting in a high effective sample size value (ESS=1829, >100). In total, 45002 trees were sampled after discharging a burn-in fraction of 25% and verifying for LnL stationarity. As convergence diagnostic, we confirmed an average standard deviation of split frequencies below 0.01 (=0.0085), and PSRFs (Potential Scale Reduction Factors) close to 1.0 (Ronquist *et al.*, 2012). Final trees were visualized with FigTree v.1.4.4 (Rambaut, 2018, http://tree.bio.ed.ac.uk/software/figtree). MEGA11 (Tamura *et al.*, 2021) was used to calculate the Kimura 2-P distance model with 1000 bootstrap and Gamma Distributed rates (Kimura, 1980).

Database sequences

- On total 123 ascidian sequences were mined from NCBI in August 2022 to use species
- delimitation analyses. Sequences were chosen following the criteria of: having the same genus
- name, given a voucher record, approved by a taxonomist, and having at least 500 bp length.
- Yet, few exceptions with sequence lengths between 450 to 500 bp were included in the analyses
- (like *Botryllus tyreus*-DQ365851- 455bp). The IDs of database sequences were given on Fig.
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Species delimitation analyses

- 142 Analyses were carried out using the Automatic Barcode Gap Discovery method (ASAP;
- Puillandre et al., 2021), a sequence similarity clustering method, and the Poisson Tree
- Processes (PTP; Zhang *et al.*, 2013), a tree-based coalescence method. The hypothetical species
- identified by these methods were assigned as Operational Taxonomic Units (OTUs). ASAP
- 146 analyses were performed on the web-based interface
- 147 (https://bioinfo.mnhn.fr/abi/public/asap/asapweb.html; accessed date: August 2022). Two
- 148 metric options provided by ASAP were used for the pairwise distance calculations; Jukes-
- 149 Cantor (JC69; Jukes and Cantor, 1969) and Kimura 2 parameter (K80; Kimura, 1980). PTP
- analyses were performed using the Bayesian implementation (bPTP), available on the web-
- based interface (http://speci es.h-its.org/ptp/, access date: August 2022). MrBayes output (tree)
- were used in the analyses by performing the default parameter values.

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Results In total, 27 ascidian sequences from Turkiye were delineated, together with 123 NCBI-mined 155 sequences. According to the common results of ASAP and PTP analyses, in total, 30 OTUs 156 were defined for the genera Botryllus, Botrylloides and Symplegma (bOTU), 1 OTU for 158 Polyclinum (pOTU) and 39 OTUs for Didemnum samples (dOTU, Fig. 2 a-c). Present study samples were located in the OTUs of 7 known species, and one sample was positioned with 160 two Saudi Arabia samples as a new Botrylloides species (Fig. 2 a,b,c). The morphological and genetic analysis results of each species are detailed below and the ASAP, PTP and LnL scores are found in Fig. S1-S3. Whereas there were common results for most of the OTUs, ASAP 162 assigned less OTU than the PTP in some groups. According to PTP analysis, almost all the D. 163 perlucidum samples were assigned to different OTUs, same results were recorded for the B. schlosseri samples. The latest OTUs were decided according to all the analysis common 165 166 results. Botrylloides niger/aff. leachii (Ascidiacea: Stolidobranchia: Styelidae: Botrylloides). In total, 221 colony fragments were collected from 6 sampling sites (Table S1), and the sequence 168 of the common haplotype (H1) was used for species delimitation analysis (Temiz et al., 2022). 170 Colonies revealed a wide range of color morphs, from a very light color (creamy) to total black (Table S2). The zooids were arranged in 'leachii type' systems (Brunetti, 2009), each ca. 1.8 172 mm long with globular stomachs and same-plane common cloacal opening, where brachial 173 sacs were occasionally seen through it (Fig. 3a₁-a₄). Two long (numbers 2-6), six middle 174 (numbers 1, 3, 4, 5, 7 and 8), and many short tentacles containing numerous blood cells fringed 175 the oral siphons (Fig. 3a₄). The present study samples were assigned in bOTU-1 together with B. niger/nigrum and B. aff. leachii, 19% distance from the closest OTUs (bOTU-6 B. leachii, 176 Fig. 2a, Table S3). 177 **Botrylloides israeliense.** In total, 36 colony fragments with light to dark color morphs (Table 178 179 S2) were collected from only Mersin sites (Mezitli, Tisan, and Kızkalesi, Table S1), and the 180 sequences of the one common (Kızkalesi sample) and one unique haplotype (Tisan sample) were used for species delimitation analysis. Zooids' lengths were ca. 2.3 mm, all arranged in a 182 'leachii type' system (Fig. 3b₁). The stomach had a globular shape continuing after a long 183 esophagus (Fig. 3b₂). The zooids were positioned vertically to ampullas (Fig. 3b₁, b₃). Four 184 long (numbers 2, 4, 6, and 8) and four medium tentacles (numbers 1, 3, 5, and 7) fringed the oral siphons (Fig. 3b₄). 185 186 The sequences of two haplotype (Tisan and Kızkalesi) were used for species delimitation 187 analysis, then were assigned to bOTU-13 with Botrylloides israeliense sample from the 188 Mediterranean coast of Israel (NCBI no:MG009580; Reem et al., 2017) and two Botrylloides 189 sp. samples (MN076468, MK978805, Viard et al., 2019) from the English Channel and the 190 Mediterranean Sea, respectively (Fig. 2a). The intra-bOTU-13 distances were 2 to 8%, and the 191 closest OTUs distanced 16 to 19% (bOTU-12, Botrylloides sp.) (Fig. 2a, Table S4). 192 Botryllus schlosseri. Four colonies collected under two rocks from the Hatay-Konacık region (Table S1), revealed three haplotypes (L12, L13 and L14; Fig. 2a). Haplotype L12 was further 193 cultured in the IMS-METU mariculture system (38-40 ppt salinity and 24-26 °C), subcloned 194 195 and colonial ramets survived under *in situ* conditions around three years. The colony's zooids 196 lengths were ca. 1.2 mm revealing a brown color morph with yellow stripes (Fig. 4a₁, 4a₂). In 197 time the pigmentation increased, and the blastogenic cycles became irregular. 198 All three Turkish haplotypes were assigned into bOTU-16 as Clade A colonies together with a database sample (MK575752). Other B. schlosseri samples mined from GeneBank were 199 200 assigned to four different OTUs; bOTU-17, -18, -19, and -20 (Table S5). The intra-bOTU-16 201 distance varied between 2 and 5%, and the highest distance was noted between our L14 sample 202 and Clade E (bOTU-20) at 19.5 % (Fig. 2a, Table S5). 203 Botrylloides anceps. In total 27 colonial samples were collected from Hatay-Konacık, Mezitli-204 Mersin, and Alanya-Antalya regions (Fig. 1, Table S1) and the sequence of the one common 205 haplotype was used for species delimitation analysis. Zooids (between 1.5-2 mm long, Fig. 4 206 and S4) were organized in 'leachii type' systems and color morphs vary from very light pink 207 to dark brown, and creamy to purple (Fig. 4b₁, b₃). Two long, four middle, and six short size 208 tentacles fringed the oral siphons (Fig. 4b₂). The common cloacal opening is quite large, 209 serving around 14 zooids (Fig. 4b₄). Stomachs with globular shapes (Fig. 4b₅). Laboratory 210 follow-up observations revealed sexual reproduction under 38-40 ppt salinity and 26-28 °C, 211 where eggs were recorded in zooids and primary buds, and testes on zooids (Fig. 4b₆). While 212 most progenies died at the oozooid stage, some survived 2-3 years under the IMS-METU 213 mariculture conditions (16 to 32 °C), while young colonies grew, without signs for 214 morphological hibernation (sensu Hyams et al., 2017, 2022), and 2-3 years old colonies

- showed exaggerated, yet condensed ampullae and irregular blastogenic cycles (all resembling
- onset of hibernation), a few months before their death (Fig. S5).
- All 27 B. anceps samples belong to a single haplotype, and the Hatay region sample sequence
- was assigned to bOTU-21 together with the Israeli *B. anceps* (MG009581, Reem *et al.*, 2017)
- and Botrylloides sp. from Puerto Rico (MT637977; Streit et al., 2021), differentiating with less
- 220 than 0.01%. Botrylloides cf. anceps (MT873573, Salonna et al., 2021) was assigned to another
- 221 OTU (bOTU-22), ca. 18 % distance from bOTU-21 (Fig. 2a, Table S6).
- 222 Botrylloides sp. A single colony (brownish with a yellow stripe around the oral siphon and
- dorsal site) of 'leachii type' systems (Fig. 5a₁) was collected from the Antalya-Kemer region
- 224 (Table S1). A colony fragment (25 zooids) was cultured for one month, revealing 1.7 mm long
- zooids (Fig. 5a₂, a₃) and blastogenic cycles of five days long under 26°C and 38-40 ppt salinity
- 226 (Fig. S6).
- 227 This colony was assigned to bOTU-8 together with two database samples from Saudi Arabia,
- Yanbu (Botrylloides sp.; ON053355-ON053356). The closest OTU (bOTU-9, Botrylloides cf.
- pannosum, MT873575; Bari, Italy; Salonna et al., 2021) is of a 14% distance (Fig 2a, Table
- 230 S7).
- 231 Symplegma brakenhielmi (Ascidiacea; Stolidobranchia; Styelidae; Symplegma): A single
- red colony was collected from the Mersin-Mezitli region under a rock (Fig. 1, Table S1). A
- ramet cultured in the aquaculture room (38-40 ppt, 27-28 °C; Fig. 5b₁) revealed transparent
- tunic, with dispersed zooids arranged with no apparent systems. Both palleal and vascular
- budding were recorded, with the latter more prominent, 1 cm long ampullae and 1-3 cm long
- 236 adult zooids (Fig. $5b_1$ - b_2).
- This specimen was assigned to bOTU-30 as S. brakenhielmi together with a Panama sample
- 238 (MT232734) at a 3% distance (Table S8, Fig. 2a) and two *Botrylloides chevalense* samples
- 239 (KX650765 and KX650764) from India, yet with a 4-5% distance (Table S8, Fig. 2a). Other S.
- brakenhielmi samples, four Indian (KU360802, MH729325, MH729324, KT276228) and one
- 241 from Italy (Bari; LS992554) samples were assigned to a different OTU (bOTU-27), together
- with S. rubra samples, ca. 28% distance to bOTU-30 (Fig. 2a, Table S8).
- 243 Polyclinum constellatum (Ascidiacea: Aplousobranchia: Polyclinidae: Polyclinum): Sixteen
- 244 colonies (light brown, reddish, and gray with a beige system of zooids) were collected only
- 245 Kızkalesi site in 2014 (Fig. 1, Fig. 5c₁).

- 246 These samples belonged to two haplotypes and were assigned into a single OTU (pOTU-1)
- 247 together with three database samples of *P. constellatum* from Italy and Florida (OL597608;
- 248 MT873559; MW285146), and with *P. indicum* and *Polyclinum sp.* (KX650781; KX650778;
- 249 KX138512) from India, with 6% intra-pOTU-1 distance (Table S9, Fig. 2b).
- 250 Didemnum perlucidum (Ascidiacea: Aplousobranchia: Didemnidae: Didemnum): This
- species was commonly observed on hard substrates of all sites (white, orange, light and dark
- creamy, and brownish color morphs). Colonies were less than 1mm thickness (Fig. 1; Table
- 253 S1; Fig. 5d₁, Table S2) and with star-shaped spicules (Fig. 5d₂).
- 254 COI sequences from two colonies (from Alanya) were used for species delimitation analysis,
- 255 together with the GeneBank data. In total, 39 OTUs were assigned for all *Didemnum* species
- 256 (Fig. 2c). According to ASAP score present study samples were clustered under the dOTU-1
- 257 together with D. perlucidum (Fig2c, Table S10) and D. etiolum (KY741541) samples from
- India, Australia and Puerto Rico with between the 0% and 5.7% distances (Table S10).

Discussion

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- This study reveals the existence and spatial distribution of eight colonial ascidian species from
- the shallow Turkish North Eastern Mediterranean coastlines. With the current sampling effort
- 263 (focused on the very shallow waters), there is a high possibility that not all colonial species
- were sampled due to seasonality or deeper ascidian habitats. In total, five botryllid ascidians
- and one Symplegma, one Polyclinum, and one Didemnum species were recorded from six sites
- between Hatay and Antalya coastlines. Distance matrix, Bayesian tree, and PTPs and ASAP
- analyses scores were used for species assignment. Whereas more OTUs were recorded in PTP
- 268 than the ASAP, common OTUs were chosen and differences were connected to data set size
- that was also experienced in the previous study (Goulpeau *et al.*, 2022).
- 270 Botrylloides niger/aff. leachii represents one of the emerged ambiguities in the taxonomic
- assignment and species delineation in botryllid ascidians (Reem et al., 2018). The first record
- of Botrylloides niger Herdman, 1886 for the Mediterranean Sea was given by Peres (1958),
- 273 and its wide distribution was reported from the Atlantic, Pacific, and Indian Ocean,
- 274 Mediterranean Sea, and the Red Sea
- 275 (https://www.marinespecies.org/aphia.php?p=taxdetails&id=252289). Based on mtCOI
- sequences, recent studies (Goutman et al., 2020; Temiz et al., 2022) reassigned the dwarf form

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of Botrylloides leachii from the Levant (Brunetti, 2009) as Botrylloides nigrum. A study (Reem et al., 2018) employing of three additional markers (18S, 28S, H3) contradicted literature suggestions, revealing minute distances between Botrylloides leachii and the 'dwarf Botrylloides leachii'. As this issue has not yet been solved, we assigned this species here as Botrylloides niger/aff. leachi. Colonies of this species were collected from all the sampling sites, representing diverse color morphotypes previously recorded in the Mediterranean Sea (Rinkevich et al., 1993; Brunetti, 2009), the Pacific and Atlantic oceans (Goodbody, 2000, 2003; Monniot & Monniot 1994). Botrylloides israeliense is a newly identified colonial ascidian (Brunetti, 2009) from the Mediterranean coast of Israel (Bay of Akko and Mikhmoret) that was already studied in the past for some ecological and biological characteristics (Rinkevich et al., 1993,1994). The genetic distances between the Turkish and the Israeli samples were between 3 to 5%, a value that is above the general eukaryotic species threshold (<2%, Hebert et al., 2004) but still lower than some other cases (~20% Resch et al., 2014; Reem et al., 2022). B. schlosseri is a highly invasive colonial ascidian distributed worldwide primarily in temperate shallow waters. Whereas the presence of the species on the Turkish coasts was reported before (Çınar, 2014; Kayış, 2011), no DNA sequence data was provided. So far, five clades have been assigned (A-E) worldwide (Reem et al., 2022), and the present study samples clustered into the most common clade (A) with up to a 5.2% intra-clade distance as compared to the worldwide within clade A distance up to 6.3%. Further, according to ASAP and pPTP results clades B, C, D, and E were assigned as different species with min.11.2% and max. 17.6% distance scores, a result that casts a query about the validity of current delimitation analyses. Botrylloides anceps is also believed to be a NIS in the Mediterranean Sea. The first record was given from the Israeli coasts by Brunetti (2009). We recorded the species for the first time in 2018 at Hatay, Mezitli, and Alanya sites. A previous visit to the sites had been done in 2012 and no single B. anceps individuals were recorded, suggesting a recent introduction by the low (0.7%) distance between the Israeli and the Turkish samples. One important morphological characteristic of this study is the variable sizes assigned to B. anceps zooids, a morphometric quality that is changed according to the colony size, adding difficulties to this simple taxonomic characteristic for ascidian species identifications.

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Colonies of presumably a new species Botrylloides sp. were recorded from only Antalya-Kemer sampling locations. This species matched 99% the NCBI samples uploaded from Saudi Arabia and the distance between the closest Botrylloides relatives was assigned at 14% (Botrylloides cf. pannosum; Australia). While this new botryllid ascidian species may have originated from the Red Sea, it has not been recorded from the Mediterranean coasts of Israel and the east sites of the Turkiye (Hatay and Mersin), thus making its origin fuzzy. The first record of Symplegma brakenhielmi in the Mediterranean Sea was from the Lebanese coast, then from the Turkish Levantine coast (Bitar & Kouli-Bitar, 2001; Çınar et al., 2006), based on only morphological characteristics. Other records included the central part of the Mediterranean Sea (Ulman et al., 2017; Mastrototaro et al., 2019). Our study on the Northeastern Mediterranean Sea reveals, for the first time, a species delimitation analysis that further indicated a possible misidentification and sequencing errors in the database, as the Turkish colonies were clustered with Panamian S. brakenhielmi and Indian B. chevalense samples, and were distinct from the Mediterranean (Bari, Italy) S. brakenhielmi cluster. Moreover, the present study sample is located as the most ancestral lineage on the tree with 100% bootstrap support which may indicate polyphyly for the genus. On the other hand, like previous study (Mastrototaro et al., 2019) S. brakenhielmi and S. rubra clustered together under the same OTU (bOTU-27), which is refer to the necessity of further morphological and genetic analysis for the whole genus. With its wide range distribution (Western Atlantic, Western Indian Ocean, Indo-Pacific, Mediterranean and Aegean), Polyclinum constellatum has been considered a cryptogenic species (Dias et al., 2013; Halim and Messeih, 2016; Aydın-Önen, 2018; Montesanto et al., 2022). Recording another name (P. indicum) under the P. constellatum OTU (pOTU1) was referred to as an error or synonymy (Montesanto et al., 2022). Present study samples were highly matched (99-100%) with database specimens uploaded from various locations all over the world (Italy, Florida, India, and the Ionian Sea), a result indicating low intraspecific distance in a widely spread species and also monophyly. Didemnum perlucidum is also a widespread species (Atlantic, Pacific, Indian Oceans, and Mediterranean Sea), and its native range is unknown (Lambert, 2002; Muñoz et al., 2015; Novak & Shenkar, 2020). Here we provide the first barcode data for the Northeastern Mediterranean Sea populations and a high-resolution phylogenetic tree, revealing genetic

distances (0 to 6%) between the widely disturbed populations (Turkiye, India, Australia, Porto-Rico). Further, whereas the Turkish samples are clustered in a single OTU together with other Didemnum perlucidum samples, other valid species were assigned in the same OTU (like dOTU4, Didemnum vexillum, and Didemnum granulatum), while samples of other inclusive species i (e.g., Didemnum psammatode) were further clustered under different OTUs (like dOTU 5, 7, and 8). All above highlighted the needs for taxonomic revision. In conclusion, we provide here the descriptions for eight colonial ascidians from the Turkish North Eastern Mediterranean Sea shallow waters, including barcode data and species delimitation, supported with major morphological features of studied species. Besides high variation in morphotypes (B. niger/aff. leachii and B. israeliense), high (B. schlosseri) and low (Didemnum perlucidum and P. constellatum) genetic diversities were also recorded. A possible new Botrylloides species (Botrylloides sp.) from the Antalya region was revealed. It is also understood that the zooid length cannot always be used as an informative taxonomic feature for B. anceps. From a phylogenetic perspective, Polyclinum members were found closer to the Styelidae members than Didemnidae and a possible polyphyly was recorded for the *Symplegma* genus and Didemnidae family.

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Figures and legends

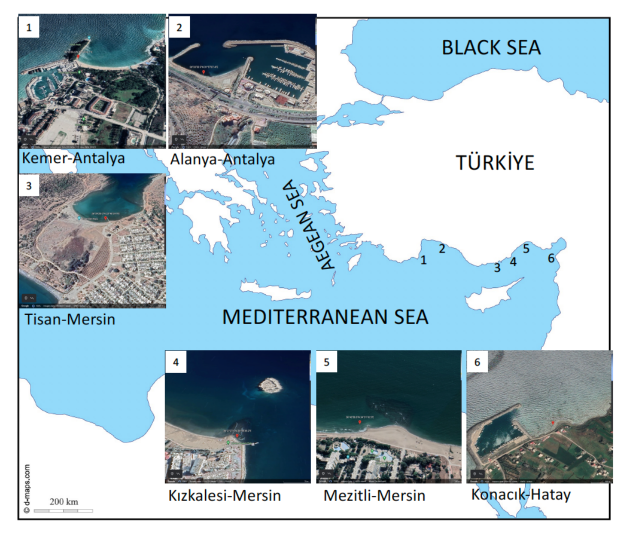


Figure 1. Map and Google Earth captures of the sampling sites. Credits; d-maps.com.

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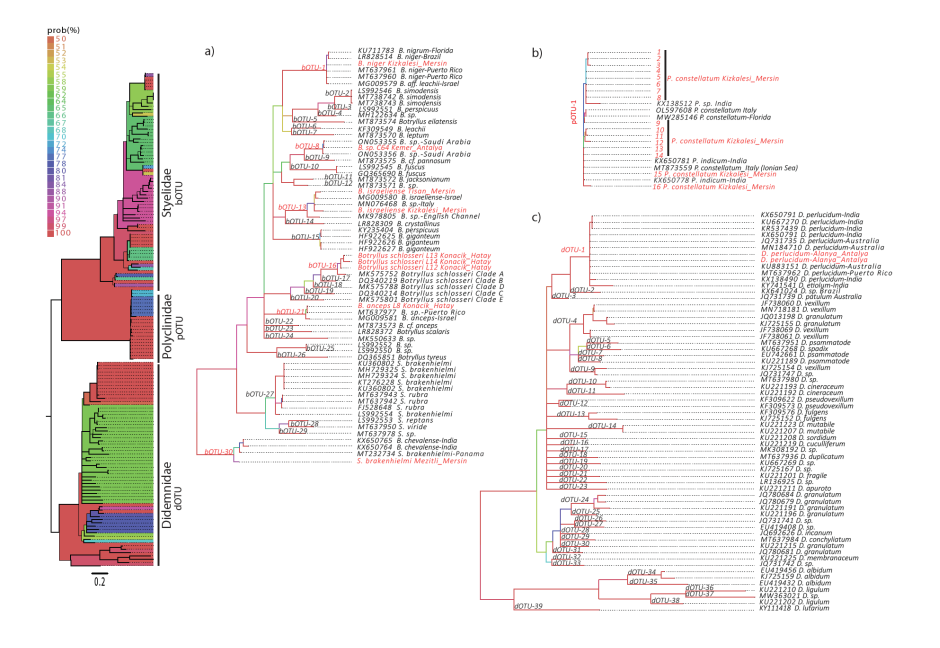


Figure 2. Bayesian majority rule consensus tree reconstructed from the ~ 600 bp COI sequence alignment. a) Styelidae specimens; b) *Polyclinum* specimens; c) Didemnidae specimens. The distance scale is given under the bootstrap probability colored tree, the NCBI IDs are provided next to the species name. OTU numbers are added on the line, according to common results of ASAP and bPTP analyses. Colors of Prob (%) scale, the main tree and a), b) and c) threes' branches represent probability values. The red colored letters are indicating the present study samples and OTUs. Abbreviations; B.: *Botrylloides*, bOTU: Styelidae family OTUs, pOTU: Polyclinidae family OTUs, dOTU: Didemnidae family OTUs.

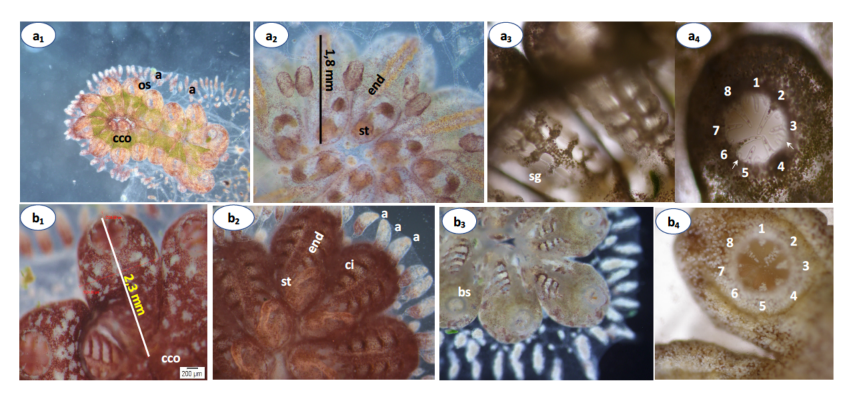


Figure 3. a₁-a₄) *Botrylloides niger/aff. leachii*, a₁) Dorsal and, a₂) Ventral view of yellow-green color morph of the colony, a₃) View of brachial sacs through the common cloacal opening, a₄) Oral siphon and tentacles; 2 long (2 and 6) and 6 middle lengths are numbered, and other short and random tentacles are indicated by arrows. b₁- b₄) *Botrylloides israeliense* b₁) Dorsal view of a zooid, b₂) Ventral view of the zooids, b₃) Dorsal view of zooids with brachial sacs (white color morph), b₄) Numbered oral tentacles (4 long and 4 medium lengths). Abbreviations: end; endostyle, ci: cell island, st; stomach, cco; common cloacal opening, a: ampullas, os: oral siphon

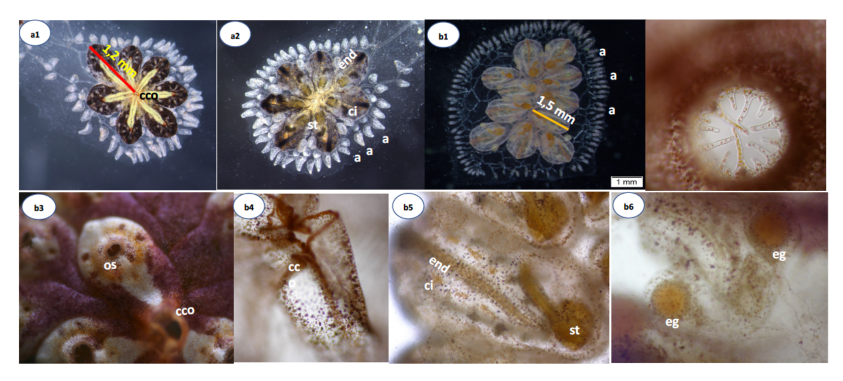


Figure 4. a₁- a₂) *Botryllus schlosseri* a₁) Dorsal and a₂) Ventral view of a cultured colony. b₁-b₆) *Botrylloides anceps*: b₁) Ventral view of a pink color morph b₂) Oral siphon and tentacles (6 long, 6 short and few very short), b₃) Dorsal view of a purple color morph zooid b₄) Common cloacal opening (cco), b₅) Single zooid ventral view, b₆) Primary bud with eggs. Abbreviations: end; endostyle, ci: cell island, st; stomach, eg; egg, cco; common cloacal opening, a: ampullas, bv; blood vessels, vb; vascular bud, os: oral siphon, co; cloacal opening.

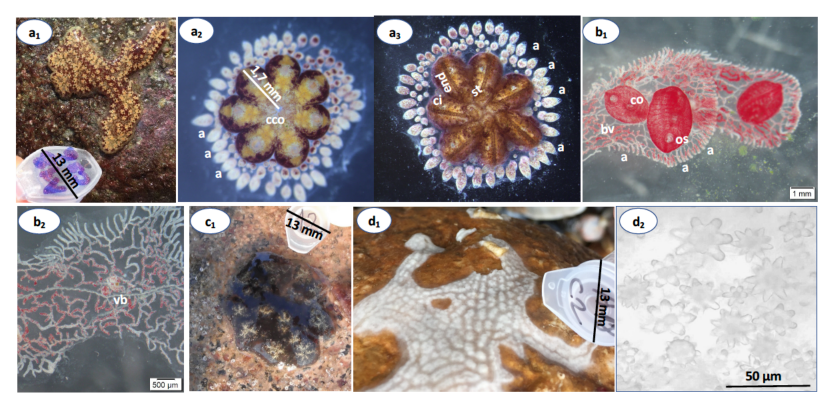


Figure 5. a₁- a₃) *Botrylloides sp.* a₁) field view a₂) cultured colony's dorsal and a₃) ventral view. b₁-b₂) *Symplegma brakenhielmi*; b₁) Dorsal and ventral views of a colony, b₂) Blood vessels and palleal budding, c₁) *Polyclinum constellatum* dorsal view, d₁-d₂) *Didemnum perlucidum*; d₁) Dorsal view of a colony, d₂) the spicules of the colony. Abbreviations: end; endostyle, ci: cell island, st; stomach, eg; egg, cco; common cloacal opening, a: ampullas, bv; blood vessels, vb; vascular bud, os: oral siphon, co; cloacal opening.

Supplementary Materials

- Table S1. Sampling sites and date details
- Table S2. General information of BoLD uploaded and other specimens
- Table S3. Kimura-2 Parameter distance for bOTU-1 and -7.
- Table S4. Kimura-2 Parameter distance for bOTU-13 and -12.
- Table S5. Kimura-2 Parameter distance for bOTU-16, -17, -18, -19 and -20.
- Table S6. Kimura-2 Parameter distance for bOTU-21 and -22
- Table S7. Kimura-2 Parameter distance for bOTU-8 and -9.
- Table S8. Kimura-2 Parameter distance for bOTU-27 and -30.
- Table S9. Kimura-2 Parameter distance for pOTU-1
- Table S10. Kimura-2 Parameter distance for dOTU-1
- Fig. S1. ASAP score; Colors represent different OTUs. Number in line of the OTUs presents the total samples assigned to the same OTU. The first number line above the OTUs' columns presents the total OTU numbers, values at the second line indicate ASAP-scores (the lowest the score the better is the partition, Puillandre et al. 2021).
- Fig. S2. PTP score; Blue lines present different OTUs, red lines present same OTUs
- Fig. S3. LnL value of MrBayes analysis.
- Fig. S4. When the zooid size of the brown morphotypes of *B. anceps* reached up to 2 mm.
- Fig. S5. Large, retracted and condensed ampullas of *Botrylloides anceps*
- Fig. S6. Blastogenic cycle of *Botrylloides sp*. Dorsal and ventral view of the Blastogenic stages; a-b) stage-D (Jun 07, 2018), c-d) stage-A (Jun 08, 2018), e-f) stage-B (Jun 09, 2018), g) Dorsal-stage C (Jun 10, 2018), h) Dorsal stage D (Jun 11, 2018). Abbreviations: end; endostyle, st; stomach, cco; common cloacal opening, sg: stigmata, a: ampullas.

Supplementary Materials: Ten supplementary tables and six supplementary figures are provided as supporting information.

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Data Availability Statement: Sequences, trace files, image files, and the primers information for each COI haplotype were uploaded to the Barcode of Life Data System (Ratnasingham & Hebert, 2007). All the data will be public upon acceptance.

Conflicts of Interest: The authors declare no conflict of interest.

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