The Origin of Asexual Brine Shrimps

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Abstract: Determining how and how often asexual lineages emerge within sexual species is central to our understanding of sex-asex transitions and the long-term maintenance of sex. Asexuality can arise “by transmission” from an existing asexual lineage to a new one through different types of crosses. The occurrence of these crosses, cryptic sex, variations in ploidy, and recombination within asexuals greatly complicates the study of sex-asex transitions, as they preclude the use of standard phylogenetic methods and genetic distance metrics. In this study we show how to overcome these challenges by developing new approaches to investigate the origin of the various asexual lineages of the brine shrimp <i>Artemia parthenogenetica</i>. We use a large sample of asexuals, including all known polyploids, and their sexual relatives. We combine flow cytometry with mitochondrial and nuclear DNA data. We develop new genetic distance measures and methods to compare various scenarios describing the origin of the different lineages. We find that all diploid and polyploid <i>A. parthenogenetica</i> likely arose within the past 80,000 years through successive and nested hybridization events that involved backcrosses with different sexual species. All <i>A. parthenogenetica</i> have the same common ancestor and therefore likely carry the same asexuality gene(s) and reproduce by automixis. These findings radically change our view of sex-asex transitions in this group and show the importance of considering scenarios of asexuality by transmission. The methods developed are applicable to many other asexual taxa.

Keywords: polyploidy, automixis, hybridization, contagious asexuality, parthenogenesis, genetic distance.

Introduction

Understanding why sexual reproduction is so widespread among eukaryotes, despite the well-known costs of sex (Maynard Smith 1978; Otto and Lenormand 2002; Meirmans et al. 2012), requires an understanding of how and how often sex-asex transitions can occur (Delmotte et al. 2001; Simon et al. 2003; Archetti 2004; Lenormand et al. 2016; Engelstädter 2017; Haag et al. 2017; Boyer et al. 2021). Here, we refer to asexual reproduction as reproduction without syngamy (i.e., without the fusion of male and female gametes but with the possibility of recombination). Most theoretical models on the origin and frequency of sex-asex transitions adopt a simplistic view and assume that an asexual clone can emerge immediately from a sexual ancestor. Although it is true that extant parthenogenetic species are derived from sexual ancestors, most of these transitions likely occurred in several steps and include nonclonal modes of asexual reproduction, which can impact the fitness of asexual lineages at both short and long evolutionary timescales (Asher 1970; Suomalainen et al. 1987; Archetti 2004, 2010; Engelstädter 2017). Historically, cytologists distinguished autotomic and apomictic asexuals according to the presence or absence of mixis (i.e., fusion of meiotic products). From a genetic standpoint, apomixis refers to clonal reproduction, which is functionally equivalent to mitosis. Cytologic and genetic definitions are contradictory because mixis is not always required for nonclonal reproduction (e.g., when either meiosis I or meiosis II is aborted; Asher 1970). Here, we use the genetic definition and refer to automixis as any modification of meiosis that leads to nonclonal inheritance (i.e., maintenance of nonzero meiotic recombination).

In this study we focus on sex-asex transitions in animals, aiming at a better understanding of both the origin of
parthenogenetic lineages and their genomic evolution. We
distinguish among four different types of origin of asexual-
ity. Asexuality can arise spontaneously (e.g., by mutation,
spontaneous origin; Simon et al. 2003), through the pres-
ence of endosymbionts (symbiotic origin; Simon et al
2003), through hybridization between two different sexual
species (hybrid origin; Cuellar 1987; Moritz et al. 1989;
Simon et al. 2003; Kearney 2005), or through transmis-
ion from an existing asexual lineage to a new one (“origin
by transmission”). Such transmission events may occur
through rare males produced by asexual lineages that may
transmit asexuality genes by mating with related sexual fe-
males (“contagious asexuality”; Hebert and Crease 1983;
Simon et al. 2003; Paland et al. 2005; Jaquiey et al. 2014).
Yet transmission through crosses between asexual females
and sexual males is also possible: either asexual females
may rarely produce reduced eggs by meiosis and undergo
rare sex or their unreduced eggs may be fertilized, leading
to an elevated ploidy level in the new lineage (e.g., produc-
tion of a triploid lineage through the fertilization, by a hap-
lloid sperm, of an unreduced diploid egg produced by an
asexual female).

Because of this diversity of possible scenarios, many
simple methods fail to provide a robust approach to eluci-
date how different asexual lineages emerge and how they
relate to each other. We can identify five major hurdles that
need to be addressed for a comprehensive understanding of
sex-asex transitions in animals. We emphasize that most of
these hurdles also undermine our understanding of sex-asex
transitions in other eukaryotes, although they may involve
other specific issues (e.g., van Dijk 2009; Lee et al. 2010).

First, although traditional phylogenetic methods can be
used to study the maternal origin of asexual lineages using
mitochondrial markers, they can be misleading in many
cases. Technically, the presence of nuclear mitochondrial
pseudogenes (“numts”; Lopez et al. 1994) can result in in-
correct inferences regarding the age of asexual lineages (Bi
and Bogart 2010). More fundamentally, classical phyloge-
netic methods based on nuclear markers might work when
asexuality arises spontaneously and when recombination is
absent, but they can be very misleading otherwise. Phylo-
genetic trees cannot depict the potentially reticulated his-
tory of asexual lineages when the origin of asexual lineages
involves crosses and/or when recombination is present. A
better approach consists in using the discordances between
mitochondrial and nuclear markers to reveal the history of
hybrid crosses (Schurko et al. 2009). Finally, with asexual-
ity by transmission, the age of asexual lineages becomes an
ambiguous concept. Indeed, different parts of the genome
of these asexuels may have experienced asexuality for very
different periods of time (Tucker et al. 2013).

The second hurdle to understanding sex-asex transitions
is that recombination may persist in asexual lineages. While
the absence of recombination in apomicts maintains het-
erozygosity levels across generations (except for mutation,
gene conversion, and mitotic recombination events), the
presence of recombination under automixis can result in
a loss of heterozygosity (LOH) between generations. When
the rate of LOH is heterogeneous across genomic regions
along the chromosome (ranging from 0% to 100%, de-
pending on the distance of the region from the centromere;
Nougué et al. 2015b; Svendsen et al. 2015; Boyer et al.
2021), different genomic regions will coalesce at different
points in time, providing different phylogenetic signals.
Hence, there are considerable uncertainties in the patterns
of molecular variation to be expected within and among
asexual genomes, as classical genetic distance metrics do
not account for heterogeneity in LOH among markers.
In addition, LOH could have different fitness consequences,
depending on the mode of asexuality (Archetti 2004, 2010;
Engelstädter 2017) or on the sex determination system of the
ancestral sexual species (Engelstädter 2008). For instance,
LOH in ZW asexual females could produce low-fitness
ZZ and WW offspring, reducing the rate of transition to
asexuality in ZW sexual species.

The third hurdle to understanding sex-asex transitions
is that asexuality is often associated with polyplody, at
least in animals (Moritz et al. 1989; Dufresne and Hebert
1994; Otto and Whitton 2000), so that studying the origin
of asexuality (e.g., allo- or autopolyplroid origin) requires
the use of specific genetic distance metrics that are defined
along the chromosome (ranging from 0% to 100%, de-
pending on the mode of asexuality (Archetti 2004, 2010;
Engelstädter 2017) or on the sex determination system of the
ancestral sexual species (Engelstädter 2008). For instance,
LOH in ZW asexual females could produce low-fitness
ZZ and WW offspring, reducing the rate of transition to
asexuality in ZW sexual species.

The fourth hurdle to understanding sex-asex transitions
is the potential occurrence of rare sex events in asexual taxa
in circumstances other than those of contagious asexuality
mentioned above. Meiosis might sometimes occur normally
in asexual females (e.g., De Meester et al. 2004; Rey et al.
2013; Boyer et al. 2021). A haploid egg produced through
regular meioses can be fertilized by a haploid sperm from
a rare male of the same or a different asexual lineage or
from a male from related sexual species. These rare events
of cryptic sex may be difficult to detect in the field, espe-
cially if divergence between parents is low or if sampling
is incomplete.

A fifth hurdle to understanding sex-asex transitions is
the technical and methodological difficulty of identifying
and exhaustively sampling the closest extant sexual species
of asexual lineages, as they often have very different geographical distributions (Kearney 2005). In addition, the closest sexual populations might be extinct or might themselves result from the hybridization between divergent sexual populations, so that many different sex-asex transition scenarios need to be considered. Overall, except in species where asexuality is directly caused by endosymbionts, ruling out a potential hybrid origin of asexuality and demonstrating that asexuality arose spontaneously remain very difficult because of this sampling challenge.

These biological and methodological challenges might exist for virtually all asexual taxa, and failing to address them consistently and jointly might provide an incomplete picture of the origin and evolution of asexual lineages. Some of these issues have recently been addressed in several systems; for instance, the description of automixis in Daphnia (Svendsen et al. 2015), the identification asexuality genes in various asexual arthropods (Sandrock and Vorburger 2011; Tucker et al. 2013; Yagound et al. 2020), and the occurrence of recombination in ancient asexuals, such as bdelloid rotifers (Simion et al. 2021).

In this study we address all of the hurdles that compromise our understanding of sex-asex transitions in Artemia parthenogenetica, a group of asexual crustaceans that has been studied for decades and is emblematic of the multiple challenges encountered when studying the origin of asexual species. This group includes diploid and polyploid asexual lineages that are found worldwide except on North America (Bowen et al. 1978; Browne 1992). The distribution of asexual lineages within the genus is exceptionally asymmetric, as they are all more closely related to Old World sexual species: Artemia sinica, A. tibetiana, A. urmiana, and an uncharacterized species from Kazakhstan (hereafter, Asin, Atib, Aurm, and Akaz, respectively; Asem et al. 2016; see table 1 for the nomenclature and abbreviations of the different taxa). The geographical distribution of both diploid and polyploid asexuals is much larger than that of these sexual species. Artemia resting stages can be dispersed by waterbirds (Sánchez et al. 2012) or humans (Rode et al. 2013) over large geographical distances.

Although the origin of asexual Artemia lineages has been extensively studied since Bowen et al. (1978), previous studies have failed to jointly address the five hurdles to understanding sex-asex transitions mentioned above. In particular, the paternal origins of asexual lineages have never been investigated, so that neither a potential hybrid origin nor a contagious origin of asexuality has been tested. Figure 1 presents a synthesis of the major conclusions of landmark articles. Briefly, asexuality in Artemia was first thought to have a single origin (Bowen and Sterling 1978; Abreu-Grobois and Beadmore 1982; Beadmore and Abreu-Grobois 1983; Abreu-Grobois 1987), such that the species was considered an “ancient asexual scandal” (Judson and Normark 1996). When both nuclear and mitochondrial markers and more sexual species (Aurm, Atib, and Asin) were considered, the evidence pointed to multiple and much more recent origins of asexuality (Baxevanis et al. 2006; Muñoz et al. 2010; Maniatsi et al. 2011; Eimanifar et al. 2015; Asem et al. 2016). These phylogenetic investigations reported that Artemia parthenogenetica was likely to be a polyphyletic group (Baxevanis et al. 2006; Maniatsi et al. 2011). According to mitochondrial data, Ap2n falls into two distinct maternal lineages, Ap2n-urm and A. parthenogenetica, whose mitochondrial haplotypes are closer to those of Akaz (mt-2nk) and Aurm (mt-2nu), respectively (Muñoz et al. 2010). A third Ap2n mitochondrial lineage has also been recently described (Maccari et al. 2013a). According to mitochondrial and nuclear data, Ap3n is thought to maternally derive either from Aurm (Maniatsi et al. 2011) or from Ap2n (Asem et al. 2016). Ap4n and Ap5n are thought to have emerged successively: the former from an Asin female, and the latter from an Ap4n female (Maniatsi et al. 2011; Asem et al. 2016).

The reproductive mode of A. parthenogenetica has also been intensely investigated for more than a century. Cytogenetic studies in diploids have yielded contradictory results by reporting almost all of the known forms of automixis (Narbel-Hofstetter 1964; Nougué et al. 2015b), while most cytogenetic studies in polyploids have reported that they reproduce through apomixis (Barigozzi 1974). Recent genetic studies showed that Ap2n reproduces through a mechanism that is genetically equivalent to central fusion automixis (Nougué et al. 2015b; Boyer et al. 2021), which leads to LOH in centromere-distal chromosomal regions owing to recombination (Svendsen et al. 2015). Artemia parthenogenetica has a ZW sex determination system and produces males at low frequency (Stefani 1964). These rare males are thought to arise through rare recombination

### Table 1: List of species names and abbreviations

| Species name          | Species abbreviation | mt abbreviation |
|-----------------------|----------------------|-----------------|
| Artemia sinica        | Asin                 | mt-sin          |
| A. urmiana            | Aurm                 | mt-urm          |
| A. tibetiana          | Atib                 | mt-tib          |
| A. sp. Kazakhstan     | Akaz                 | mt-kaz          |
| Diploid A. parthenogenetica (2n) with Aurm-type mitochondria | Ap2n-urm | mt-2nu |
| Diploid A. parthenogenetica (2n) with Akaz-type mitochondria | Ap2n-kaz | mt-2nk |
| Tripliod A. parthenogenetica (3n) | Ap3n | mt-3n |
| Tetraploid A. parthenogenetica (4n) | Ap4n | mt-4n |
| Pentaploid A. parthenogenetica (5n) | Ap5n | mt-5n |
events between the Z and W chromosomes in Ap2n females, which results in a LOH at the sex locus and the production of ZZ males (Browne and Hoopes 1990; Abreu-Grobois and Beardmore 2001; Boyer et al. 2021). Last, for a long time rare males have been thought to be useless and irrelevant to Ap2n reproduction (MacDonald and Browne 1987; Simon et al. 2003), but experiments showed that they can transmit asexuality when crossed with Aurm and Akaz sexual females (Maccari et al. 2014; Boyer et al. 2021). Moreover, recent experiments have shown that some Ap2n females can engage very rarely in sex in the laboratory, with normal meiosis and recombination (Boyer et al. 2021). Hence, contagious asexuality and/or sexual reproduction may occur at small rates within Ap2n asexuals in nature.

In addition, previous studies also suffered from a number of limitations. The first limitation is the lack of reliable nuclear markers available across asexual and sexual taxa. The most informative data set (23 allozyme markers) dates back to the 1980s but shows a limited resolution with few alleles per marker (Abreu-Grobois and Beardmore 1982). Similarly, nuclear sequences (ITS1, Na+/K+-ATPase, etc.) show limited diversity across asexual and sexual taxa (Baxevanis et al. 2006; Asem et al. 2016). Microsatellite markers described in Muñoz et al. (2008) present null alleles and fail to amplify in some sexual species (Maccari et al. 2013b). Mitochondrial data also present limitations due to the potential coamplification of nuclear pseudogenes (Wang et al. 2008). Indeed, several studies have reported difficulties in amplifying cytochrome c oxidase subunit I (COI) sequences using universal primers or primers designed for the distantly related Artemia franciscana (Wang et al. 2008) and A. salina (Asem et al. 2016). Mitochondrial-nuclear (hereafter, “mitonuclear”) discordance represents crucial evidence for hybridization scenarios, yet many studies did not combine nuclear and mitochondrial data (fig. 1) and relied only on mitochondrial data for taxonomic identification (Sainz-Escudero et al. 2021). Furthermore, in many studies ploidy of all samples is not directly assessed or is based on existing literature regarding populations previously sampled in the same locality (e.g., Baxevanis et al. 2006). However, a same locality may host different (sexual and/or asexual) populations whose occurrence varies spatially or temporally (e.g., Agh et al. 2007). Importantly,
ploidy cannot be assessed according to the number of alleles at each locus, as _Artemia_ individuals from each ploidy level often exhibit only two alleles at each locus (Nougé et al. 2015a); whether this lack of variation is the result of LOH or of some other mechanisms has never been studied. Finally, no previous study has included all known sexual species and all ploidy levels. A failure to include the most closely related sexual species may result in asexuality appearing more ancient than it actually is (e.g., Perez et al. 1994). Similar biases due to limited sampling are frequent in studies of the age and origin of asexual taxa (Tucker et al. 2013).

In this article we investigate potential origins by transmission of diploid and polyplloid asexual lineages, considering all new experimental information regarding the reproductive mode of _A. parthenogenetica_ (Maccari et al. 2014; Nougé et al. 2015b; Boyer et al. 2021). We test the hypothesis that diploid and polyplloid asexual lineages may represent a mixture of different “hybrids” resulting from several events of contagion and secondary backcrosses with different sexual species. We build a series of tailored population genetic methods to test whether asexual _Artemia_ of various ploidies have a hybrid origin, and we attempt to identify the corresponding parental species. We also investigate whether secondary crosses or contagion events can be identified. To this end, we conduct the first study that includes an exhaustive sampling of asexual lineages (Ap2n, Ap2n-urm, Ap3n, Ap4n, Ap5n) and major sexual relatives (Aurim, Akaz, Atib, Asin) and that combines nuclear and mitochondrial data with ploidy data (based on flow cytometry). Finally, we also test for the presence of cryptic sex in Ap2n. In the absence of sexual reproduction, we can indeed expect Ap2n individuals with different mitochondrial haplotypes to be characterized by different and specific nuclear backgrounds. In contrast, in the presence of cryptic sex, we expect a discordance between mitochondrial haplotypes and nuclear backgrounds. We find that after a single hybrid origin of one diploid asexual lineage, all other asexuals emerged through a series of four nested hybridization events involving several sexual species. Overall, this new approach changes our view of asex-asex transitions in _Artemia_. It may prove to be a valuable tool to investigate sex-asex transitions in other taxa, especially when classical phylogenetic approaches are not appropriate.

Methods

Samples

On the basis of the existing literature (Abreu-Grobois and Beardmore 1982; Muñoz et al. 2010; Maniatsi et al. 2011; Maccari et al. 2013a), we chose 37 populations from Eurasia and Africa, including both asexual strains (described as diploid, triploid, tetraploid, and pentaploid) and the four closest sexual species (fig. 2). We obtained samples from cyst bank collections and wild-collected adults (fig. 2; table S4; tables S1–S5 are available online). Cysts were hatched and individuals were maintained following protocols described in Rode et al. (2011). It has recently been shown that at least some Ap2n females can reproduce sexually in the laboratory at a rate of ~2% in the presence of males (Boyer et al. 2021). However, for simplicity (and because the capacity to undergo rare sexual reproduction in _natura_ is unknown), we categorized individuals only as sexuals or asexuals and did not consider facultative asexuality. The reproductive mode of each population was verified according to the presence or absence of males among adults. When at least one male was present, we separated asexual females from sexual females according to morphological characters (Maccari et al. 2013b). All five populations with at least one male consisted of a mixture of asexual females with females from different sexual species (Artemia franciscana: AIM, BOL, SAG; A. sinica: DON; or A. salina: BDP).

Ploidy

We characterized the ploidy of putatively diploid and polyplloid asexual females as well as of males and females of each sexual species (147 individuals total) using flow cytometry as described in Nougé et al. (2015b) and section 1 of the supplemental PDF. Data from Nougé et al. (2015b) were added to the data set, resulting in a total sample size of 206 individuals (table S4). We tested for difference in genome size between asexual and sexual lineages using t-tests. All significant P values remained significant after Bonferroni correction, so we present only the uncorrected values for simplicity.

COI Genotyping

A fragment of the mitochondrial COI gene of 336 individuals (table S4) was amplified using primers 1/2COI_Fol-F and 1/2COI_Fol-R following the protocol of Muñoz et al. (2010). Because these primers turned out to lack specificity and resulted in the amplification of a large number of numts (see below), we amplified the COI sequence of 23 additional individuals using the more specific primers Co1APAR-F(5'-TTTGGAGCTTGAGCAGGAAT-3') and Co1APAR-R(5'-TGCGGGATCAAAGAAAGAAG-3'; see the supplemental PDF, sec. 2). Polymerase chain reaction (PCR) products were purified and directly sequenced (i.e., with no cloning) using an ABI PRISM BigDye Terminator kit (Applied Biosystems, Warrington, United Kingdom) on an ABI PRISM 3130xl sequencer. After removing
**Figure 2:** Geographic distribution of the sexual and asexual *Artemia* samples genotyped in this study. Because of the scale of the map, the ANK (Ankiembe, Madagascar) *A*. *p3n* population is represented with an arrow pointing toward its location. Mitochondrial data could not be obtained for KOY (Koyashskoye, Ukraine), which is represented according to the analysis of 17 *A*. *p2n-kaz* individuals from Maccari et al. (2013a) that used the same cyst sample. Each color corresponds to a mitogroup. AIB = Aibi Lake (China); AIM = Aigues-Mortes (France); ANK = Ankiembe (Madagascar); ARS = Arcos de la salinas (Spain); ATA1/2 = Atanasovko Lake (Bulgaria); BAM = unknown location, Bameng area (China); BDP = Bras del Port (Spain); BET = Bethioua (Algeria); BOL = Bolshoye Yarovoe (Russia); BUJ = Bujaraloz (Spain); CIT = Citros (Greece); COQ = Co Qen (China); DON = Dongiagou (China); IMO = Imón (Spain); IZM = Izmir (Turkey); KAZ = unknown location (Kazakhstan); KUL = Kulundinskoye (Russia); LAG = Lagkor Co (China); LAM = La Mata (Spain); LAR = Larache (Morocco); LAV = Lavalduc (France); LPM = La Palmes (France); MAH = Maharlu Lake (Iran); MOL = Molentargius (Italy); NAR = Narte (Albania); ODI = Odiel (Spain); ROC = Nuestra Señora del Rocío (Spain); SAG = Salin De Giraud (France); TEK = Teke Lake (Kazakhstan); TEN = Tene (Spain); TNG = Tanggu (China); URM = Urmia Lake (Iran); VIL = Sète-Villeroy (France); XIE = Xiechi Lake (China); YIM = unknown location, Yimeng area (China); YIN = Yingkou (China).
sequences with indels and sequences from the same individual that were identical, we recovered 359 sequences that were split into two data sets: data set 1, which included only high-quality sequences (198 sequences including at least 538 sites in the final alignment and without any ambiguous position), and data set 2, which included all other sequences (161 sequences that were either too short or included one or more ambiguous positions). COI phylogenetic inferences are often biased by the inclusion of numts (Buhay 2009) or chimeric sequences that result from the coamplification by PCR of mitochondrial sequences with contaminating sequences (either numts or mitochondrial sequences from other individuals; e.g., Dubey et al. 2009). We developed a new method to identify potential chimeric sequences based on a data set composed of known numts sequences (see “Analyses of Mitochondrial Data 3: Chimera Detection” below). To build this data set, we performed additional cloning and sequencing of PCR fragments amplified either from total DNA or from DNA enriched from mitochondrial DNA (supplemental PDF, sec. 2; table S4). Cloning allowed the recovery of 32 numt sequences without indels (which we identified as minority sequences among those obtained from the same individual through cloning (supplemental PDF, sec. 2) and seven mitochondrial sequences (hereafter, data set 3). We built a data set that combined these 39 sequences (data set 3), our 198 sequences (data set 1), and 748 Artemia spp. high-quality sequences from GenBank without any indel (supplemental PDF, sec. 2). All analyses of the final data set of 985 COI sequences were performed in R version 3.6.3 (https://www.r-project.org).

Analyses of Mitochondrial Data 1: Pseudogene Detection

We aligned sequences using MAFFT (ver. 7.427; Katoh et al. 2002) as implemented within the package ips (ver. 0.0.11; Heibl 2008) with default settings. To detect potential numts without indels, we translated sequences into amino acid sequences using the invertebrate mitochondrial DNA genetic code. To detect potential numts, we tested for changes in the polarity of amino acid residues (Kunz et al. 2019). For each sample, we estimated the absolute difference in polarity (i.e., PP1 in Cruciani et al. 2004) between derived and ancestral amino acid sequences. On the basis of the observed distribution of this polarity difference across the 985 protein sequences (fig. S1; figs. S1–S3 are available online; supplemental PDF, sec. 3), we set 0.1 as the threshold in polarity differences above which sequences were labeled as potential numts. This procedure allowed the reliable detection of 65% of the reference numts obtained by cloning (i.e., 21 of 32 known numts). We also detected 26 additional sequences (including 25 sequences from GenBank) that were labeled as potential numts for the rest of the analyses.

Analyses of Mitochondrial Data 2: Haplotype Reference Set

Poorly aligned positions were removed using Gblocks (ver. 0.91b) as implemented within the package ips (ver. 0.0.11; Heibl 2008). They were collapsed into 230 unique haplotypes using the haplotype function of the package haplotypes (ver. 1.1.2; Aktas 2020).

Analyses of Mitochondrial Data 3: Chimera Detection

We designed a quantitative test to detect and exclude potential chimeric sequences. The principle of the method is to determine, for each focal sequence, how many mutations could be “explained” by assuming that this sequence represents a chimera. We compared each haplotype sequence with the remaining 229 sequences in the data set to identify the most similar one. For each mutation differing between the focal sequence and the most similar one, we provide a score of one if this mutation (i.e., the same single-nucleotide polymorphism [SNP]) was found in another unrelated sequence of the data set or zero otherwise. For each of the 230 haplotype sequences, we computed the sum of this score over the different SNPs of each haplotype. We considered that each sequence could have acquired a mutation that happened to be present in an unrelated sequence of the data set (score = 1) but that it could not have acquired two or more of these mutations (score > 1). Hence, haplotype sequences with a score equal to or greater than two were considered potentially chimeric (for details, see the supplemental PDF, sec. 4). We removed 80 haplotypes (corresponding to 101 samples) that appeared as potential chimeras resulting from the coamplification of a mitochondrial sequence and either a numt or contamination from other mitochondrial sequences. Overall, we obtained 123 reference non-chimeric haplotypes (corresponding to 884 samples).

Analyses of Mitochondrial Data 4: Haplotype Assignment

To study the phylogenetic relationship among major asexual and sexual taxa (Ap2n-kaz, Ap2n-urm, Ap3n, Ap4n, Ap5n, Aurm, Akaz, Atib, Asin), we built a haplotype network based on the 123 reference haplotypes and using the parsimnet function (95% probability of parsimony; Templeton et al. 1992) of the package haplotypes (ver. 1.1.2; Aktas 2020). We found the same tree topology when building a maximum likelihood phylogeny with the phangorn package (ver. 2.5.5; Schliep 2011).

In addition, we assigned the 61 Ap2n sequences in data set 2 to Ap2n-kaz, Ap2n-urm, or other sequences (i.e., numts or chimeras). To do so, we aligned them with the 123 reference haplotype sequences and with the 107 numt and chimeric sequences and estimated pairwise genetic
distances as described above. For each sequence in data set 2, the mitochondrial haplotype was assigned according to the identity of the closest reference haplotype(s). Whenever a numt, a potential numt, or a chimera was found among the closest sequences, the Ap2n mitochondrial haplotype was set as “unknown.” We could successfully assign 46 of the Ap2n sequences to either Ap2n-kaz or Ap2n-urm.

**Microsatellite Genotyping**

We genotyped 432 individuals with a panel of 12 microsatellite markers (for details regarding markers and amplification protocol, see Muñoz et al. 2008; Nougué et al. 2015a). Data from Nougué et al. (2015b) were added to the data set, resulting in 489 typed individuals (table S4; supplemental PDF, sec. 5). Standardization was achieved by adding DNA from the same individual onto the different plates. Genotype data at three microsatellite markers (Apdq01TAIL, Apdq02TAIL, Apdq03TAIL) were used only when investigating the relationship among Ap2n but excluded when analyzing the full data set because of the presence of null alleles in polyploids and sexuals (Maniatsi et al. 2011).

**Microsatellite Data Analyses 1: Lynch Distance**

The Lynch genetic distance is a genetic distance estimate based on a band-sharing index (i.e., one minus the similarity index, eq. [1] in Lynch [1990]). It is an appropriate metric to broadly compare well-separated sexual and asexual groups with different ploidy levels, such as Ap2n, Ap3n, Ap4n, Ap5n, and the different sexual species (i.e., ignoring the occurrence of LOH in Ap2n). We computed the average pairwise nuclear distance between and within lineages using the Lynch genetic distance, as implemented in the polysat package (ver. 1.7-4; Clark and Jasieniuk 2011) in R. To visualize distance data, we transformed individual pairwise distances into principal coordinate axes using the cmdscale function in the stats package (ver. 3.6.3). To investigate the homogeneity of the reference sexual populations, we investigate whether they show a signal of admixture (Estoup et al. 2016). We examined the variation in allele sizes within individuals from each sexual taxon (Aurm, Akaz, Atib, Asin; supplemental PDF, sec. 6).

**Microsatellite Data Analyses 2: Genetic Distance for Automictic Ap2n**

Because it does not account for the possibility of LOH, the Lynch genetic distance is of limited use regarding relationships within Ap2n. We therefore present a new genetic distance measure for automicts, accounting for the different possible paths between diploid genotypes (supplemental PDF, sec. 6; table S1). We assumed that mutations rate is the same across loci but that LOH varies across loci proportionally to the average inbreeding coefficient (Fis), which can be independently estimated. This new measure weights events according to the relative magnitude of LOH and mutation events (with rates r and µ, respectively). This genetic distance is a proxy for the time length of the path between individuals (or averaged across different possible paths, according to their relative probability of occurrence). For instance, with this new distance measure, two individuals with genotypes AA and AB are weighted as distant if the locus has a strongly negative Fis (low LOH rate), since their difference likely results from a single A-to-B mutation. However, they are not weighted as distant if the locus has a strongly positive Fis (high LOH), as AA can result from a LOH event from an AB parent.

**Monophyly of Ap2n-kaz and Ap2n-urm Clades**

In the absence of sexual reproduction, we expect Ap2n-kaz and Ap2n-urm individuals to be characterized by different and specific nuclear backgrounds. In contrast, in the presence of sexual reproduction between Ap2n-kaz and Ap2n-urm individuals, we expect mitonuclear discordance. Using data from the 12 microsatellite markers, we used our new genetic distance metric to compute pairwise genetic distances among 127 Ap2n individuals with known mitochondrial haplotypes. Using a randomization test, we first investigated whether the genetic distance between Ap2n-kaz and Ap2n-urm lineages was significantly larger than that within Ap2n-kaz and Ap2n-urm lineages. As this test considers only average nuclear distances between and within lineages, the difference might be significant even if Ap2n-kaz and Ap2n-urm clades are not monophyletic (e.g., because of rare events of sexual reproduction between Ap2n-kaz and Ap2n-urm individuals). Using the pairwise distance matrix computed above, we first built a neighbor-joining (NJ) tree using the nj function from the ape package (ver. 5.4-1; Paradis and Schliep 2019) and estimated branch length using nonnegative least squares (nnsle.tree function from the phangorn package ver. 2.5.5; Schliep 2011). We computed the 95% confidence interval of this branch length by resampling microsatellite markers to build 1,000 bootstrap replicates. Finally, we built NJ trees separately for Ap2n-kaz and Ap2n-urm and assembled them into a single tree where each lineage is monophyletic. We again estimated the branch length of this tree using nonnegative least squares and then tested whether the estimated branch length was outside of the 95% confidence interval computed above.

**Evolutionary Origin of Ap2n-kaz and Ap2n-urm**

To investigate the evolutionary relationships between Ap2n-kaz and Ap2n-urm and the three sexual species
(Aurm, Akaz, and Asin), we considered different scenarios based on the Lynch genetic distance (fig. 3). We considered two independent spontaneous origins within Aurm and Akaz (fig. 3A), a single spontaneous origin followed by a hybridization event (fig. 3B), two independent hybridization events (fig. 3C), and one hybridization and one backcross event (fig. 3D). These four scenarios were considered with or without the presence of an unknown species, denoted Aunk, which is assumed to carry mt-urm. We did not consider the scenarios where Aunk carried mt-kaz. Indeed, nuclear data indicated that all Ap2n are much closer to Akaz than to Aurm (see “Results”), and it is therefore very likely that Ap2n-kaz inherited their mitochondria directly from Akaz. Because the out-group Asin can have different positions (it can be closest to Akaz, Aurm, or Aunk), each scenario was evaluated assuming all three possible topologies. For each topology, we described each branch length by a parameter. The number of identifiable parameters is given for each scenario in figure 3. For scenarios in which Ap2n arose spontaneously, a branch length between Ap2n and the sexual species was included. When Ap2n arose from a hybrid cross, the genetic distance between them and a given sexual species was computed as the averages of the branch lengths between either parent and that sexual species. The model corresponding to each scenario was fitted using least squares to the matrix of Lynch genetic distance among Akaz, Aurm, Asin, Ap2n-kaz, and Ap2n-urm. To avoid any confounding effect due to potential cryptic sex between Ap2n-kaz and Ap2n-urm, we computed this matrix after removing 13 Ap2n individuals with discordant mitonuclear data (see “Results”). We assumed that genetic distance within each species or within each of two Ap2n lineages was negligible compared with the genetic distance among species (i.e., we ignored divergence within each sexual or asexual lineage). Models were compared according to the corrected Akaike’s information criterion (AICc; Hurvich and Tsai 1989). We computed the difference (ΔAICc) between the AICc of a given model and that of the model with lowest AICc. Models with ΔAICc higher than two were considered poorly supported (Burnham and Anderson 2002).

Figure 3: Scenarios for the origin of Ap2n-kaz and Ap2n-urm. The two lineages are depicted by light orange and dark orange circles, respectively. The color of the dot within each circle represents the maternal lineage of the mitochondrion (blue indicates mt-2n from an Aurm mother, and yellow indicates mt-2nk from an Akaz mother). White dotted lines and black dashed lines indicate spontaneous and hybrid origins, respectively. In A and in the presence of Aunk, the nine branch lengths are not identifiable. The branch length leading to Aunk cannot be fitted and was therefore dropped. In B, the scenarios illustrated assume a spontaneous origin of Ap2n-kaz in Akaz followed by a cross with Aurm or Aunk. We also considered the reciprocal scenario (origin in Aurm or in Aunk and cross with Akaz). In D, the scenarios illustrated show an origin of Ap2n-urm through hybridization followed by a backcross with Akaz. The reciprocal scenario with a backcross with Aurm or Aunk was also considered. Only one topology (where Asin is closest to Aunk) is represented for all scenarios involving Aunk (bottom row). The two other topologies were also considered (Asin closest either to Akaz or Aurm). The best model is indicated by the red rectangle and involved one hybridization with an unknown species and backcross with Akaz. The number of parameters fitted for each model (corresponding to the number of identifiable branch lengths) is given by the number in parentheses below each scenario.
Evolutionary Origin of Ap3n, Ap4n, and Ap5n

We assumed the maternal origin of the ancestor of Ap3n, Ap4n, and Ap5n lineages to be known according to the mitochondrial data (see "Results"). For each ploidy level, we compared different scenarios involving different paternal origins (for details, see the supplemental PDF, sec. 7). For each ploidy level and each scenario, we simulated 10,000 synthetic hybrids using a custom script in R. For each hybrid, we first randomly sampled a mother with the observed mitochondrial haplotype in our data set (i.e., Ap2n-kaz for Ap3n, Asin for Ap4n, and Ap4n for Ap5n). Second, to draw a haploid genotype (representing the sperm genotype), we randomly sampled an individual for each paternal origin (i.e., Akaz, Ap2n-kaz, Ap2n-urm, Aurm, Atib, or Asin) and randomly sampled one allele of this individual at each locus. We assumed that the 12 microsatellite loci were unlinked, so that probabilities of sampling alleles were independent across loci. We assumed that alternative scenarios involving fertilization by an unreduced sperm were less likely (e.g., origin of Ap3n through the fertilization of a reduced Ap2n-kaz egg by an unreduced sperm or origin of Ap4n through the fertilization of an unreduced Asin egg by an unreduced sperm). We then computed the average Lynch genetic distance based on the 100 synthetic hybrids closest to Ap3n, Ap4n, or Ap5n individuals in our data set.

Results

Ploidy Characterization

The results from the flow cytometry measurements are summarized in figure 4. The ploidy levels detected in each population were in good agreement with those found in previous cytological or genetic studies (Abreu-Grobois and Beardmore 1982; Muñoz et al. 2010; Maccari et al. 2013a), except for two Ap5n populations previously described as Ap4n (BUJ and CIT; Abatzopoulos et al. 1986; Amat et al. 1994; Maniatis et al. 2011). Interestingly, the genome size of Ap2n-kaz was not significantly different from that of Ap2n-urm (Ap2n-kaz: 4.74 pg, SD = 0.17; Ap2n-urm: 4.65 pg, SD = 0.20; $t = -1.93$, df = 50.48, $P = .059$). The Ap2n-kaz genome size was significantly lower than Akaz (4.92 pg, SD = 0.14; $t = -3.21$, df = 14.74, $P = .006$), and the Ap2n-urm genome size was significantly higher than Aurm (4.22 pg, SD = 0.20; $t = 4.46$, df = 5.67, $P = .005$). The genome size of Ap3n (7.13 pg, SD = 0.43) was consistent with their ploidy level and was not significantly different from 1.5 times that of Ap2n-kaz ($t = -0.16$, df = 37.63, $P = .88$). Although Ap4n harbor an Asin mitochondrial (Asem et al. 2016), their genome size (10.15 pg, SD = 0.34) was more than twice that of Asin (2 $\times$ 4.74 = 9.49 pg, SD = 0.24; $t = 4.44$, df = 6.38, $P = .004$). In contrast, the size of the genome of Ap5n (12.22 pg, SD = 0.45) that also harbor an Asin mitochondrion was not significantly different from 2.5 times that of Asin ($t = 2.15$, df = 4.57, $P = .09$). These observations seem consistent with the scenario of an origin of Ap4n through an endoduplication in Asin. Interestingly, the genome size of Atib was 57%, 34%, and 39% larger than that of Aurm, Akaz and Asin, respectively, suggesting an increase in genome size in the lineage leading to Atib.

COI Genotyping

Ap2n, Ap3n, Ap4n, and Ap5n were found in 24, 3, 7, and 5 populations, respectively (table S4; fig. 2). Among the 37 populations sampled, only five (ATA, IZM, NAR, COQ, and DON) were composed of individuals with different ploidies. Similarly, among the 24 Ap2n populations, 13 comprised only individuals with Ap2n-kaz haplotypes, five comprised only individuals with Ap2n-urm haplotypes, and six (AIM, ATA, VII, SAG, IZM, NAR) comprised individuals with both Ap2n-kaz and Ap2n-urm haplotypes (fig. 2). Consistent with previous studies, Ap2n-kaz is the clade with the largest geographic distribution (fig. 2). When including all sequences found in GenBank, the distribution of Ap2n-urm is restricted to the Mediterranean area (Spain, France, Italy, and Turkey), around the Black Sea (Atanasovoko Lake, Bulgaria; Oybuskoye Lake, Ukraine), and in western China (Aibi Lake, Lagkor Co). Polyploids also have a large geographic distribution (fig. 2).

Among the 950 sequences from data set 1 and from GenBank, we found 47 sequences that had a large change in the polarity of the amino acid sequence and 88 sequences that included two or more mutations found in another unrelated sequence of the data set. These sequences were likely to be mutants and chimeras, respectively. The remaining 850 COI sequences were collapsed into 123 unique haplotypes of diploid mt-2n; diploid mt-2n; triploid, tetraploid, and pentaploid asexual lineages; and related sexual species (fig. 5). In line with Maccari et al. (2013a), we found three networks separated by more than 30 mutation steps. The fourth network described in their study corresponded to pseudogenes (GenBank accession: EF615587-8) that differ from other sequences by several nonsynonymous mutations that changed polarity. Each asexual lineage was characterized by a majority haplotype found at high frequency in many populations and recently derived satellite haplotypes found at a lower frequency in a few populations (fig. 5), as previously observed in Ap2n lineages (Muñoz et al. 2010; Maccari et al. 2013a). This observation of low haplotypic diversity with a starlike shape is consistent with a recent range expansion of the different asexual lineages, which now have widespread geographical distributions. No haplotype was shared between asexual lineages and either of the sexual
Figure 4: Estimated genome size of Artemia sexual and asexual lineages. Genome size (pg) of diploid Ap2n-kaz and Ap2n-urm; triploid, tetraploid, and pentaploid asexual lineages; and related sexual species (Aurm, Asin, Akaz, and Atib). Mean ± SD C values are shown. See figure 2 for population abbreviations. Numbers below population labels indicate sample sizes. Thirty-seven Ap2n individuals had unknown mitochondrial haplotypes and are not represented.
Figure 5: Statistical parsimony network of mitochondrial haplotypes from diploid mt-2n\(k\), diploid mt-2nu, triploid mt-3n, tetraploid mt-4n, and pentaploid mt-5n asexual lineages and related sexual species (mt-urum: Aurm; mt-sin: Asin; mt-kaz: Akaz; and mt-tib: Atib). Haplotypes including samples with different ploidies are represented with pie charts. Circle diameter is proportional to the relative haplotype frequency among the 850 sequences that were neither numts nor chimeras. Connecting lines indicate single substitutions, and small black circles represent putative missing haplotypes. Haplotype codes correspond to those reported in table S5. Footnotes are as follows. (a) KBG4 sequence of a cyst from Kara-Bogaz-Gol (Turkmenistan) was molecularly assigned to Aurm by Eimanifar et al. (2014). (b) Sequences from five Ap3n individuals (based on cytology; Asem et al. 2016) from Aibi Lake (China) had an APD01 Ap2n-kaz haplotype. (c) Sequences from five Ap2n individuals (based on cytology; Asem et al. 2016) from Akkikol Lake (China) had an ANK1 Ap3n haplotype. (d) Sequences from 52 Ap2n individuals (based on morphology; Muñoz et al. 2010; Mascari et al. 2013; Eimanifar et al. 2014, 2015) had the same APD05 haplotype as Ap3n individuals in our data set. (e) The TU13 Asin sequences from Siberia were found to be chimeric between the Ap4nARS1 mitochondrial haplotype and an Ap5n numt; it was included to illustrate that Ap4n and Ap5n likely originated from Siberian Asin.
species. The majority haplotype of Ap2n-kaz differed by six mutations from the closest Akaz haplotype (but one haplotype of Ap2n-kaz differed from Akaz by just one mutation). The majority haplotype of Ap2n-urm differed by one mutation from the closest Aurm haplotype. The majority haplotype of Ap3n was identical to the closest Ap2n-kaz haplotype, and Ap4n and Ap5n had the same majority haplotype, which differed by a single mutation from the closest Asin haplotype (fig. 5). We found an 18% divergence between Aurm and Asin COI haplotypes from reference mitochondrial genomes used in Sainz-Escudero et al. (2021), which corresponds to a divergence time of 6.6 Ma (5.47–7.07 Ma). Using divergence between Atib and Asin COI haplotypes and the divergence time estimate from Sainz-Escudero et al. (2021) provides a qualitatively similar age estimate of 0.062 Ma (0.051–0.079 Ma). Although these results rely on strong assumptions (accurate dating of the fossil used for calibration and constant molecular clock), they suggest that all extant asexual lineages potentially emerged more recently than previously thought (i.e., less than 80,000 years ago).

The first network includes haplogroups corresponding to Ap2n-kaz, Ap2n-urm, Ap3n, Aurm, Akaz, and Atib. This result confirms the existence of the two distinct major Ap2n lineages, Ap2n-kaz and Ap2n-urm. The third Ap2n lineage described in Maccari et al. (2013a) possibly represents a chimera between two divergent PCR-amplified sequences in that study (fig. S2). Consistent with Maccari et al. (2013a), the haplogroup mt-tib1 included most Atib sequences from Lagkor Co (also known as Gaize Lake; Zheng and Sun 2013).

Triploid samples (Ap3n), whose assignation is based on flow cytometry and nuclear genotype, were found to be nested within Ap2n-kaz (fig. 5). This suggests that triploids are maternally derived from this diploid lineage. Ap3n were characterized by a very low mitochondrial diversity (table 2) despite their large geographical distribution (Madagascar, Turkey, Albania). When including sequences from other studies (retrieved from the National Center for Biotechnology Information), we found triploids with sequences identical to those of the closely related Ap2n-kaz haplotypes, and we observed diploids with sequences identical to those of Ap3n haplotypes (fig. 5).

Among sexuals, Aurm had a larger haplotypic diversity than Akaz and Atib from Lagkor Co. This is likely due to a larger sample size, which increases the likelihood of sampling rare alleles (fig. 5; table 2).

A second network consisted of one Atib sequence from Lagkor Co (AtibLAG2) and sequences from other sexual populations (Haiyan Lake, Jingyu Lake, Nima, Yangnapengco, Qi Xiang Cuo, etc.) from the Qinghai-Tibet Plateau that we refer to as mt-tib2 (fig. 5). This haplogroup differed by more than 35 mutation steps from mt-tib1 and segregates at a frequency of ~3% in Lagkor Co (1 of 34 Atib sequences). No sequence from the mt-tib1 haplogroup was found in populations from Haiyan and Jingyu Lakes. Only a few sequences per population were available from GenBank, so making general conclusions is not possible.

The third network included sequences from Ap4n, Ap5n, and Asin. We found that some Ap4n and Ap5n samples shared the same mitochondrial haplotype (fig. 5). We found low haplotype diversity among Ap4n and Ap5n haplotypes (in contrast to Asem et al. 2016; table 2). The diversity found in Ap4n and Ap5n by Asem et al. (2016) was due to 12 sequences, which were likely to be numts or chimeras according to our analysis. Ap4n and Ap5n were closest to a haplotype from an Asin sample from Lake Dus-Khol (also known as Lake Svatikovo, East Siberia, Russia; Naganawa and Mura 2017), which was found to be chimeric between the Ap4n mitochondrial haplotype and an Ap5n numt (table S5). This suggests that Asin from Lake Dus-Khol, Ap4n, and Ap5n share the same mitochondrial haplotype and at

Table 2: Mitochondrial (Kimura two-parameter model on cytochrome c oxidase subunit I sequences) genetic distances between and within mitogroups of sexual species and asexual lineages

|           | Aurm  | Ap2n-urm | Ap2n-kaz | Ap3n  | Akaz  | Atib (LAG) | Atib (others)* | Asin  | Ap4n  | Ap5n  |
|-----------|-------|----------|----------|-------|-------|------------|----------------|-------|-------|-------|
| Aurm      | .007  | .009     | .026     | .030  | .018  | .017       | .075           | .184  | .206  | .206  |
| Ap2n-urm  |       | .004     | .029     | .033  | .021  | .020       | .080           | .188  | .210  | .210  |
| Ap2n-kaz  | .026  |          | .005     | .007  | .016  | .015       | .061           | .175  | .196  | .197  |
| Ap3n      | .030  | .029     |          | .003  | .021  | .015       | .083           | .180  | .202  | .202  |
| Akaz      | .018  | .033     | .021     |       | .003  | .017       | .073           | .175  | .197  | .197  |
| Atib (LAG)| .017  | .021     | .015     | .003  |       | .015       | .073           | .175  | .180  | .180  |
| Atib (others)* | .075 | .080     | .015     | .007  | .021  | .017       | .074           | .175  | .197  | .197  |
| Asin      | .184  | .080     | .083     | .073  | .017  | .021       | .074           | .175  | .197  | .197  |
| Ap4n      | .206  | .196     | .196     | .202  | .175  | .180       | .184           | .167  | .017  | .021  |
| Ap5n      | .206  | .197     | .197     | .202  | .175  | .180       | .184           | .167  | .021  | .002  |

Note: See table S5 for the accession numbers of the 123 unique haplotype sequences. See table 1 for abbreviations.
* Distances based on Atib sequences from populations other than Lagkor Co (LAG) were computed separately.
least one numt. Hence, \textit{Ap4n} might have originated in East Siberia from an \textit{Asin} mother. Mitochondrial diversity levels in the \textit{Asin} population from Xiechi Lake (\textit{AsinXIE} and \textit{XIE} haplotypes in fig. 5) were similar to those observed in other sexual species.

\textit{Microsatellite Genotyping}

Genetic distances within and among asexual and sexual lineages are represented in figure 6 and table 3. The proportion of total variability explained by the first, second, and third axes of the principal coordinate analysis were 28.6\%, 25.9\%, and 19.2\%, respectively (fig. 6). We observed a larger genetic diversity within both \textit{Ap2n-kaz} and \textit{Ap2n-urm} than within \textit{Ap3n}, \textit{Ap4n}, or \textit{Ap5n} polyploids. Sharing of multilocus genotypes among populations was rare except for some geographically close populations (\textit{Ap2n}: VII/AIM, YIN/DON; \textit{Ap5n}: IZM/ATA/CIT). Consistent with mitochondrial data, \textit{Ap2n-kaz} were more closely related with \textit{Akaz} than with \textit{Aurm}. Surprisingly, \textit{Ap2n-urm} were also more closely related to \textit{Akaz} than to \textit{Aurm}. Compared with \textit{Ap2n-kaz} and \textit{Ap2n-urm}, \textit{Ap3n} were more closely related to \textit{Aurm}. Consistent with mitochondrial data, \textit{Ap4n} and \textit{Ap5n} were more closely related to \textit{Asin} compared with \textit{Ap2n-kaz}, \textit{Ap2n-urm}, and \textit{Ap3n}. Variation in microsatellite allele size was smaller in \textit{Akaz} and \textit{Asin} (YIM population) than in \textit{Aurm} and \textit{Atib}, which suggests that the later two populations might be admixed (fig. S3; supplemental PDF, sec. 6).

\textit{Monophyly of Ap2n Mitochondrial Clades}

The NJ tree based on our genetic distance for automicts, which accounts for recombination and null alleles, is represented in figure 7. Individuals that cluster together in this tree often had the same mitochondrial haplotype group (\textit{Ap2n-kaz} or \textit{Ap2n-urm}). In other words, nuclear

\textbf{Figure 6:} Principal coordinate analyses of asexual lineages and sexual species. The first three principal coordinate axes are shown, with the percentage of variation represented by each axis given between parentheses. Sexual and asexual taxa are represented with circles and squares, respectively. Numbers within squares represent the ploidy level. \textit{Ap2n} individuals with unknown mitochondrial haplotypes are shown in gray.
genetic distance between pairs of individuals with mt-2nk and mt-2nn mitochondrial haplotypes was larger, on average, than that between pairs of individuals with the same mitochondrial haplotype group (fig. 7; for Ap2n-kaz–Ap2n-urm, distance \(d = 9.23\); for Ap2n-urm–Ap2n-kaz, \(d = 8.51\); for Ap2n-kaz–Ap2n-kaz, \(d = 8.22\)). This association between mitochondrial haplotype and nuclear background was highly significant: among 10,000 randomizations of the data set, the estimated genetic distance between pairs of individuals with the same mitochondrial haplotype was never lower than observed genetic distances of 8.51 or 8.22 (i.e., \(P < .0001\)). We identified 13 individuals with a potential mismatch between mitochondrial haplotype and nuclear background (arrows in fig. 7). Importantly, the mitochondrial haplotypes of these individuals were neither unique nor shared with sexual species (Aurm, Akaz) but rather corresponded to major Ap2n haplotypes. Moreover, according to the bootstrap analysis, the total length of the NJ tree was not significantly lower than a tree with forced monophyly of Ap2n-urm and Ap2n-kaz (i.e., without mitonuclear discordance; \(P = .295\)). Hence, we could not rule out the hypothesis that the mitonuclear discordance patterns occurred by chance.

### Evolutionary Origin of Ap2n-kaz and Ap2n-urm

To compare the different scenarios for the evolutionary origin of Ap2n-kaz and Ap2n-urm, we used the average pairwise genetic distance between Ap2n-kaz, Ap2n-urm, Akaz, Aurm, and Asin (table 3). The scenario with the emergence of Ap2n lineages through one hybridization and one backcross event (fig. 3D) had the highest support (table S2). According to this scenario, Ap2n-urm first arose through a hybridization event between an Akaz male and an Aunm female, carrying a mt-urm. Ap2n-kaz arose through a backcross between an Akaz female and an Ap2n-urm rare male. All other scenarios could be ruled out (\(\Delta A I G_c > 2.5\); table S2), including scenarios involving a spontaneous origin, two independent hybrid origins, and a different order of events (e.g., Ap2n-kaz being the F1 and Ap2n-urm being the backcross).

### Evolutionary Origin of Ap3n, Ap4n, and Ap5n

For Ap3n, average genetic distances inferred from the scenario involving an Aurm paternal origin were significantly lower than other genetic distances (\(P < 2E-16\); table S3; fig. 8A). For Ap4n, average genetic distance inferred from the scenario involving an Ap2n-kaz paternal origin was slightly better than that with an Ap2n-urm paternal origin (\(P = .029\); table S3; fig. 8B). For Ap5n, average genetic distances inferred from the scenario involving an Atilb paternal origin were significantly lower than genetic distances from other scenarios (\(P < .0003\); table S3; fig. 8C). All data underlying these results and figures have been deposited in the Dryad Digital Repository (https://doi.org/10.5061/dryad.7h4j0zsb; Rode et al. 2022).

### Discussion

#### General Implications for the Study of the Origin and Evolution of Asexuals

Our study proposes new ways to investigate the origin of asexual lineages. We try to address the five hurdles that frequently face such studies using approaches that can likely be used in asexual taxa other than *Artemia parthenogenetica*. First, when considering potential hybrid or contagious origins of asexuality, our study showcases that the genealogy of asexual groups might include several origins and/or several nested hybridization events. Hence, the origin of asexuality gene(s) should be clearly distinguished from the origin of asexual clades that carry these genes. Indeed, we find a single origin of asexuality gene(s) in *A. parthenogenetica*, followed by several nested hybridizations with different sexual species. Second, heterogeneity in LOH within and across chromosomes should be carefully considered in automictic asexuals. In particular, with central fusion automictics, which
are frequent, heterozygosity can be high close to the centromere but decreases farther away from the centromere. Thus, LOH creates a heterogeneity in coalescence times both among markers along each chromosome and between asexuality gene(s) and the rest of the genome. The methods developed in this study should improve genetic distance estimates between automictic asexual lineages by explicitly accounting for this heterogeneity. In contrast to a widespread hypothesis regarding the evolution of asexuality in ZW lineages (Engelstädter 2008), we show that this sex...
Figure 8: Comparison of the different scenarios for the paternal origin of Ap3n (A), Ap4n (B), and Ap5n (C). For each ploidy level and each paternal origin, we estimated Lynch genetic distance across nine microsatellite loci between 10,000 simulated hybrids and real individuals of our data set with the corresponding ploidy level (Ap3n: n = 16; Ap4n: n = 23; Ap5n: n = 15). The horizontal line and the red dot represent the median and the mean, respectively, of the 100 synthetic hybrids with the smallest genetic distance. See text for details.
determination system does not represent a major obstacle to the evolution of asexuality, provided that recombination is low enough (or quickly evolved to low levels; Boyer et al. 2021). In addition, in these systems the production of rare ZZ males may allow for the production of new asexual lineages through contagious asexuality. Third, our results support the hypothesis that polyploidy is often a consequence rather than a cause of asexuality (Neiman et al. 2014). In addition, our findings suggest that asexual groups with odd ploidy levels (3n, 5n) can reproduce through automixis. This implies that despite the uneven number of chromosomes, pairing between some homologous chromosomes does occur but that this pairing does not prevent the formation of an unreduced egg. Fourth, the occurrence of rare sex can greatly impact the genomic evolution of asexual lineages by increasing their diversity. The genomic consequences of these events depend on whether they occur within an asexual lineage, between different asexual lineages, or between sexual lineages and related sexual species. Rare sex events within an asexual lineage can be detected only when there is some genetic diversity within asexual populations. Importantly, selection might favor a particular asexual lineage within a population and may locally erase the genetic footprints of rare sex events, increasing the difficulty to detect them. Rare sex between different asexual lineages or between asexual lineages and related sexual species can be detected through admixture tests or through cytonuclear discordances. The new method developed here to test for cytonuclear discordances assesses whether groups defined according to mitochondria are monophyletic at the nuclear level. This method should be widely applicable in other systems. Fifth, although our sampling was as exhaustive as possible, our results point to particular geographic locations toward which sampling efforts of sexual relatives should be directed in the future to refine the estimates of the age for the origin of asexuality in Artemia (see below).

**Origin of A Sexuality in Artemia: Overview**

The most parsimonious scenario to summarize our main results is presented in Figure 9 and indicates that both diploid and polyploid asexual Artemia harbor nuclear genomes that are admixed between the nuclear genomes of Akaz and one or several of the other sexual species. Additional sampling and more genetic data may reveal that some specific crosses are actually more complex and/or that other scenarios need to be statistically compared (e.g., scenarios involving serial backcrosses).

The relatively low diversity in mitochondrial haplotypes suggests that contagious asexuality is present but rare in Artemia (or leads most of the time to unfit hybrids). Despite the relatively frequent occurrence of rare males in Ap2n (Maccari et al. 2013b), our data suggest that contagious asexuality via rare males has occurred twice, leading to the Ap2n-kaz and Ap2n-urm lineages, with no evidence for further successful events of contagious asexuality. Indeed, the diversity of mitochondrial haplotypes within each of the main asexual lineages is limited, and these haplotypes are not intermingled with haplotypes found in sexual lineages. Mutation (rather than backcrossing with unsampled sexual populations) thus seems the most parsimonious explanation for the observed haplotype diversity within each main lineage of asexual Artemia.

In addition, each mitochondrial haplotype is only a few mutational steps away from the mitochondria of the different sexual species (Akaz, Aurm, Asin), which suggests that all of the extant asexual lineages are more recent than previously thought (less than 80,000 years old compared with more than 3 and 0.84 Ma; Perez et al. 1994; Baxevanis et al. 2006; Eimanifar et al. 2015). This estimate represents the age of the most recent common ancestor of all of these lineages (i.e., the oldest extant lineage Ap2n-urm, from which all of the other lineages likely arose through subsequent hybridization events). Importantly, the loci that cause asexuality may be much older than our rough estimate of the age of their common ancestor. Indeed, we cannot exclude...
that all of the original asexual lineages went extinct, leaving only recently derived ones. This issue can be solved only by comparing the divergence of the region(s) directly associated with asexuality with that of the rest of the genome (Tucker et al. 2013). We discuss the implications of these findings for *A. parthenogenetica* asexuality before discussing each of these events in detail.

**Diploids and Polyploids May All Be Automicts**

Our findings strongly suggest that all asexual *Artemia* probably derive from a single-original hybrid ancestor. The same asexuality genes are therefore very probably shared among all diploid and polyploid *A. parthenogenetica*. We found that the apparent polyphylly of the group (Baxevanis et al. 2006) results from a history of nested crossings between asexuals and sexual relatives.

Automixis in *Ap2n* has been a major source of confusion throughout one century of cytological observations. Recent genetic data have clarified this debate and support the conclusion that *Ap2n* reproduce by central fusion automixis (Nougue et al. 2015b; Boyer et al. 2021). In contrast, most of the literature on polyploids is not controversial and claims that they are apomictic (Brauer 1894; Artom 1931; Barigozzi 1944, 1974). Reproduction of polyploids via an automictic process that would involve recombination has been ruled out on the basis of three types of observations. First, polyploids do not seem to produce rare males (Goldschmidt 1952; Metalli and Ballardin 1970; Chang et al. 2017). Second, each ploidy level shows high heterozygosity but little clonal diversity (Abreu-Grobois and Beardmore 1982; Zhang and King 1992; Maniatsi et al. 2011). Third, cytological observations have been claimed to refute automixis (because of the failure to observe meiosis, the authors inferred that sister chromatids separate through a mitosis; Barigozzi 1974).

Asexuality is likely to have the same genetic determinism in diploid and polyploid asexual lineages, since they share a common asexual ancestor. If true, this means that the distinction between *Ap2n* automicts and polyploid apomicts may be erroneous. Beyond shared ancestry, several lines of evidence support that polyploids may well, in fact, have the same reproductive mode as diploids.

First, the cytological evidence is not as clear-cut as often reported. This confusion relies on different definitions of automixis by cytologists and geneticists. The former use the fusion of meiotic products as a criterion, while the latter use the genetic consequences of modified meiosis as a criterion (Asher 1970; Nougue et al. 2015b; Svendsen et al. 2015). Most cytologists did not consider the possibility that automixis could occur through the abortion of one of the two meiotic steps. In fact, this aborted meiosis has been described by Goldschmidt (1952) in *Ap3n* and *Ap5n* polyploids. She did not observe any fusion of meiotic products but a brief

synapsis and the production of a polar body. Furthermore, she observed that the number of elements (bivalent or univalent) drops during diakinesis and increases afterward to reach univalent number at metaphase. These observations refute the occurrence of apomixis. They show that meiosis I is aborted at the end of prophase I (with bivalents being separated within the oocyte but the resulting univalents not being distributed to different daughter cells and instead realigned at the equatorial plate) and jumps directly to metaphase II. Note that in *Ap3n* and *Ap5n*, this brief pairing of homologues can easily occur despite the odd number of chromosomes (one chromosome simply stays unpaired during prophase I), as observed in other animal species (e.g., Christiansen and Reyer 2009). This meiosis modification corresponds to the reproductive mode of *Ap2n* (Nougue et al. 2015b; Boyer et al. 2021) and exactly matches “central fusion automixis” as defined by geneticists. According to Asher (1970), the suppression (or abortion) of meiosis I is genetically equivalent to automixis with central fusion as defined cytologically (i.e., where fusion actually occurs), while suppression or abortion of meiosis II is cytologically equivalent to automixis with terminal fusion. Other authors refer to the suppression (or abortion) of one of the meiotic divisions as “meiotic apomixes” (e.g., Archetti 2010). However, this term refers to both the suppression of meiosis I and the suppression of meiosis II, whose outcome differs genetically when recombination is present. For this reason, we prefer the term “central fusion automixis” in the large, genetic sense, noting that it includes meiotic apomixis with suppression of meiosis I.

Second, if polyploids are automicts, they may occasionally lose heterozygosity because of recombination, as observed in *Ap2n* lineages (Boyer et al. 2021). However, there are good reasons to expect that in polyploids, homeologous chromosomes (i.e., pairs of chromosomes derived from the two parental species of allopolyploids; Glover et al. 2016) are more divergent than nonhomeologous chromosomes (i.e., pairs of chromosomes derived from only one of the two parental species) and that they will pair much less frequently, which likely results in almost no LOH between homeologs. This also likely limits recombination-generated genetic variation in polyploids. In addition, it might also drastically reduce the rate of rare male production in polyploids. In *ZZW* triploids, *Z* chromosomes are likely to preferentially pair and recombine (leaving the *W* unpaired), before all chromosomes realign on the equatorial plate and meiosis II starts. Similarly, in *ZZWW* tetraploids or *ZZZZW* pentaploids, *W* nonhomeologous chromosomes are likely to preferentially pair together (and even if *Z* and *W* homeologous chromosomes would pair, two subsequent LOH events would be required to produce *ZZZZ* or *ZZZZZ* males). Furthermore, as in all polyploids, there is probably a strong selection pressure to reduce the number of
crossovers to avoid interlocking crossover events among different pairs of chromatids (Lenormand et al. 2016), which may reduce recombination in polyploids and reinforce this apparent apomictic-like reproduction. As we often observe only two alleles in polyploids, recombination rates between nonhomeologous chromosomes might be small but greater than the mutation rate. Hence, the absence of rare males in polyploids is not necessarily an argument against central fusion automixis. Polyploids *A. parthenogenetica* may therefore not be apomicts. They may reproduce by central fusion automixis, but polyploidy and the absence of pairing between homoeologous chromosomes would make this reproductive mode genetically very close to apomixis. This interpretation is open to further tests, but it would explain why polyploids have apomictic-like reproduction while being derived from automic lineages, as suggested by our study.

**Hybrid Origin of Ap2n-urm and Contagious Origin of Ap2n-kaz via an Ap2n-urm Rare Male**

Our best scenario is that *Ap2n-urm* likely arose through hybridization between an *Akaz* male and a female from *Aurm*, an unknown sexual species with a mitochondrial haplotype close to that of *Aurm*. It is quite likely that *Aurm* would be related to *Akaz* and *Aurm* and therefore be present in Crimea or in Central Asia, if not extinct. Interestingly, one sexual population from Crimea could be a good candidate. It is currently described as an *Aurm* population (Abatzopoulos et al. 2009; Maccari et al. 2013a), but unexpectedly, its mitochondrial haplotype is closer to that of *Ap2n-urm* than to that of *Aurm* (one vs. three mutational steps, respectively, in the network in fig. 5). Unfortunately, we could not obtain this sample for this study.

According to our best model, the second *Ap2n* group originated through a backcross between a rare male of this first lineage and an *Akaz* female. The genome size of *Ap2n-kaz* is consistent with this scenario. The backcross might have occurred almost immediately after the first occurrence of *Ap2n-urm*, as rare males may be produced at a higher rate in young asexual lineages (Boyer et al. 2021) and as both crosses rely on the presence of *Akaz*, which likely has a very limited geographical range, at least today. All *Ap2n* lineages seem to branch from these two major groups, and we did not find firm evidence of any further event of contagious asexuality. Such secondary contagion through rare males would indeed capture new mitochondrial haplotypes from sexual species, which would be easily detected. The *mPAK1* sequence in figure 4 from Maccari et al. (2013b) might be a rare candidate, but without nuclear data, we cannot further investigate this possibility. Overall, the low diversity among *Ap2n* haplotypes suggests that secondary contagion is probably very rare. The restricted geographical distributions of sexuals in Central Asia and around the Crimean Peninsula and their spatial or temporal segregation from sexuals (Mura and Nagorskaya 2005; Shadrin and Anufrieva 2012) might indeed limit the chances of contagion.

Recent experimental evidence (Boyer et al. 2021) has shown that *Ap2n-kaz* may occasionally reproduce sexually. Reproduction between *Ap2n-kaz* and *Ap2n-urm* could perhaps explain some mitonuclear discordance observed in our data (fig. 7). Alternatively, *Ap2n-kaz* and *Ap2n-urm* might form monophyletic clades, and these discordances might be due to our limited number of microsatellite markers. Finally, sexual reproduction might preferentially occur within *Ap2n-kaz* and *Ap2n-urm*, as most populations are composed of individuals from a single clade (fig. 2). Mitochondrial divergence is low within *Ap2n-kaz* and *Ap2n-urm*, so that we could not test and detect mitonuclear discordance within each clade with our method. Additional mitochondrial and nuclear data will be needed to better test these different hypotheses.

**Origin by Transmission of Ap3n via an Ap2n-kaz Female**

All *Ap3n* branch within the *Ap2n-kaz* lineage and show very limited genetic diversity at both the mitochondrial level and the nuclear level despite having a worldwide distribution, from Madagascar to the Mediterranean. This result is in line with that of Maniatsi et al. (2011) that found only two different clones across 10 *Ap3n* populations. It contrasts with the large allozyme diversity reported within and among three *Ap3n* populations by Abreu-Grobois and Beardmore (1982). Our samples (NAR, IZM, ANK) belong to a single *Ap3n* lineage that likely arose through a cross between an *Ap2n-kaz* female and an *Aurm* male. When considering samples from previously published studies, we found several mitochondrial haplotypes that are shared between *Ap3n* and *Ap2n-kaz* (fig. 5; supplemental PDF, sec. 9). This might have been caused by improper assignation of ploidy levels of these samples or by the independent origin of several highly related, but distinct, *Ap3n* lineages. Further sampling, genotyping, and careful checking of ploidy levels should resolve the matter.

**Contagious Origin of Ap4n via an Ap2n-kaz Rare Male**

We found that *Ap4n* show very limited genetic diversity, consistent with previous studies (Abreu-Grobois and Beardmore 1982; Maniatsi et al. 2011). In the most likely scenario (fig. 8), *Ap4n* resulted from a standard hybrid cross between reduced haploid gametes from an *Asin* mother and an *Ap2n-kaz* rare male (other scenarios involving unreduced *Asin* gametes are less likely, as they require the combination of multiple rare events). Tetraploidy would have occurred through endoduplication in one of these
hybrids (or within its descendants). This scenario is consistent with the observation that most microsatellite loci have only two alleles in those individuals. The genome size of Ap4n is 7% larger than that of Asin and Ap2n-kaz combined, which suggests an increase in genome size following polyploidization. It is unlikely that Ap4n have multiple origins, since we would expect to observe the capture of several mitochondrial haplotypes given the mitochondrial diversity observed in Asin. Historically, the hybridization event probably took place in East Siberia, as Ap4n carry a mitochondrion that is found only in an Asin sample from this area (for a discussion of mitochondrial diversity within Asin, see the supplemental PDF, sec. 9). Finally, all of the Ap4n lineages observed in the Mediterranean area would be a subsample of few successful lineages, which reached a worldwide distribution.

**Origin by Transmission of Ap5n via an Ap4n Female**

According to mitochondrial data (fig. 5), Ap5n are derived from an Ap4n unreduced gamete fertilized by a male of unknown origin. The microsatellite data agree with this scenario, as the Ap5n genotype is very close to Ap4n and suggests an Atib paternal origin. However, according to flow cytometry data, the genome size of a hybrid resulting from the fertilization of an Ap4n unreduced egg by a reduced Atib sperm would be 10.0% larger than the observed genome size of Ap5n. This discrepancy may be due to secondary reduction in Ap5n genome size or to an incorrect paternal assignment. The second- and third-best candidates would be Ap2n-kaz and Akaz, respectively, and the genome size of hybrids resulting from these crosses would better match that of Ap5n (2.4% and 3.1% larger, respectively). Additional nuclear data will indicate whether Atib paternal origin is the correct inference.

**Implications for the Study of the Origin of Asexual Lineages**

A more robust dating of the origin of asexuality in Artemia and establishing firm scenarios for the origin of the different asexual lineages require the different sexual species to be more extensively sampled and characterized. Ap2n-urm is clearly more closely related to the sexual Crimean population from Lake Koyashskoe than to Aurm according to both mitochondrial data (fig. 3) and nuclear data (Abatzopoulos et al. 2009; Sainz-Escudero et al. 2021). This suggests that the Lake Koyashskoe population might be Aurm. Additional nuclear genetic data from this and other populations (e.g., Kara-Bogaz-Gol) are required to confirm this hypothesis. Similarly, mitochondrial and nuclear genetic data from sexual populations from the Altai region currently described as Aurm (Mura and Nagorskaya 2005; Shadrin and Anufriieva 2012) could help decipher whether one of these populations actually corresponds to the Akaz population involved in the paternal origin of Ap2n-urm and/or the maternal origin of Ap2n-kaz. In addition, studying Asin populations from Siberia (which are very divergent from Asin populations from China; fig. 5) should shed light on the maternal origin of Ap4n. Finally, studying the diversity of Atib populations outside the reference population used to describe this species (Lagkor Co) could help assess the paternal origin of Ap5n. Indeed, we confirm that most individuals from Lagkor Co have mt-tib1 mitochondrial haplotypes very close to mt-kaz (fig. 5), whereas all other sexual populations from the Qinghai-Tibet Plateau assigned as Atib carry only the very divergent mt-tib2 haplotype (as previously found in Maccari et al. 2013a). Our microsatellite data suggest that the Lagkor Co population might be admixed (fig. S3). Hence, it is possible that the Lagkor Co population has been introgressed by a mitochondrial haplotype closely related to mt-kaz (Maccari et al. 2013a). Alternatively, all other populations may represent another undescribed sexual species that can hybridize with Atib individuals, resulting in a low frequency of the divergent mt-tib2 haplotype in Lagkor Co. Similarly, our microsatellite data suggest that Aurm might be admixed between two divergent populations (fig. S3), which is consistent with the observation of two very divergent ITS1 haplotypes that segregate within Aurm (Eimanifar et al. 2014; Sainz-Escudero et al. 2021). Additional genetic data from populations from the Qinghai-Tibet Plateau and from Crimea will be required to disentangle different admixture scenarios and resolve potential mitonuclear discordances in Atib and Aurm.

**Taxonomic Implications**

The taxonomy of asexual lineages is often ambiguous, as clearly exemplified by our case study. Provided that multiple hybridization events can give rise to different asexual lineages, phylogenetically defined monophyly is a poor criterion of the taxonomic description of asexual lineages. Asexuality also prevents the use of species delimitation criteria based on interfertility between different taxa. Because of these difficulties, it may be more biologically relevant to focus on the common origin of asexual lineages and their common reproductive mode. In our case study it makes sense to collectively refer to A. parthenogenetica as a relevant taxonomic unit, as they probably inherited the same asexuality gene(s) from their common ancestor. The drawback of this taxonomic approach, in Artemia and other groups, may be that a single name hides the diversity of hybridization events that led to the different lineages. As a consequence, a minimal convenient way to designate these taxa would be to distinguish the major groups derived from these crosses, if they can be distinguished. For instance, in A. parthenogenetica,
five major groups may summarize the major sex-asex transitions: Ap2n-kaz, Ap2n-urn, Ap3n, Ap4n, and Ap5n. Finally, taxonomic issues, lineage history, and age may also be better resolved by directly identifying and studying the genomic regions associated with asexuality (Sandrock and Vorburger 2011; Tucker et al. 2013; Yagound et al. 2020). As our case study shows, sex-asex transitions are quite different from the idealized view of an apomictic mutant arising in a sexual species. Ultimately, a better characterization and understanding of sex-asex transitions represents a pivotal step to refine our theories for the long-term maintenance of sexual reproduction and the extant distribution of sexual and asexual taxa.

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Statement of Authorship

N.O.R., F.D., and T.L. conceived the study; F.H. and G.V.S. provided samples; N.O.R. and R.J.-Z. performed flow cytometry measurements with advice from F.D.; E.F., R.J.-Z. and F.D. performed genotyping; N.O.R., L.B., C.H., and T.L. reviewed the existing literature; N.O.R. and T.L. analyzed the data and wrote the manuscript with input from the co-authors; and T.L. managed the project and funding.

Data and Code Availability

The 234 sequences from data sets 1 and 3 have been deposited in GenBank (accession numbers are provided in table S5). Microsatellite and flow cytometry data, as well as scripts used for data analysis, have been deposited in the Dryad Digital Repository (https://doi.org/10.5061/dryad.7b44j0zs; Rode et al. 2022).

Literature Cited

Abatzopoulos, T. J., F. Amat, A. D. Baxevanis, G. Belmonte, F. Hontoria, S. Maniatsi, S. Moscatello, et al. 2009. Updating geographic distribution of Artemia urmiana Günther, 1890 (Branchiopoda: Anostraca) in Europe: an integrated and interdisciplinary approach. International Review of Hydrobiology 94:560–579.

Abatzopoulos, T. J., C. D. Katrissis, and C. D. Triantaphyllidis. 1986. A study of karyotypes and heterochromatic associations in Artemia, with special reference to two N. Greek populations. Genetica 71:5–10.

Abreu-Grobois, F. 1987. A review of the genetics of Artemia. Pages 61–99 in P. Sorgeloos, D. A. Bengston, W. Declair, and B. Jaspers, eds. Artemia research and its applications. Vol. 1. Universa, Wetteren, Belgium.

Abreu-Grobois, F. A., and J. A. Beardmore. 1982. Genetic differentiation and speciation in the brine shrimp Artemia. Pages 345–376 in C. Barigozzi, ed. Mechanisms of speciation. Liss, New York.

Beardmore, J., and F. Abreu-Grobois. 1983. Taxonomy and evolution of sexual and asexual taxa. Evolutionary Biology. Springer, Boston.

Baxevanis, A. D., I. Kappas, and T. J. Abatzopoulos. 2006. Molecular phylogenetics and asexuality in the brine shrimp Artemia. Molecular Phylogenetics and Evolution 40:724–738.

Beardmore, J., and F. Abreu-Grobois. 1983. Taxonomy and evolution in the brine shrimp Artemia. Pages 153–164 in G. Oxford and D. Rollinson, eds. Protein polymorphism: adaptive and taxonomic significance. Academic Press, New York.
Bi, K., and J. P. Bogart. 2010. Time and time again: unisexual salamanders (genus *Ambystoma*) are the oldest unisexual vertebrates. BMC Evolutionary Biology 10:238–252.

Bossier, P., W. Xiaomei, F. Catania, S. Dooms, G. Van Stappen, E. Naessens, and P. Sorgeloos. 2004. An AFLP database for authentication of commercial cyst samples of the brine shrimp *Artemia* spp. (International Study on *Artemia* LXX). Aquaculture 231:93–112.

Bowen, S. T., and G. Sterling. 1978. Esterase and malate dehydrogenase isozyme polymorphisms in 15 *Artemia* populations. Comparative Biochemistry and Physiology B 61:593–595.

Browne, R. A., and C. W. Hoopes. 1990. Genotype diversity and selection in asexual brine shrimp (*Artemia*). Evolution 44:1035–1051.

Buhay, J. 2009. COI-like sequences are becoming problematic in molecular systematic and DNA barcoding studies. Journal of Crustacean Biology 29:96–110.

Burnham, K., and D. Anderson. 2002. Model selection and multi-model inference: a practical information-theoretic approach. 2nd ed. Springer, New York.

Chang, M., A. Asem, and S. Sun. 2017. The incidence of rare males in seven parthenogenetic *Artemia* (Crustacea: Anostraca) populations. Turkish Journal of Zoology 41:162–222.

Christiansen, D. G., and H.-U. Reyer. 2009. From clonal to sexual reproduction of water frogs. Evolution 63:1754–1768.

Cuculo, G., M. Baroni, E. Carosati, M. Clementi, R. Valigi, and S. Clementi. 2004. Peptide studies by means of principal properties of amino acids derived from MIF descriptors. Journal of Chemometrics 18:146–155.

Cudiar, O. 1987. The evolution of parthenogenesis: a historical perspective. Pages 43–97 in P. Moens, ed. *Mémosis*. Academic Press, Orlando, FL.

De Meester, L., A. Gómez, and J. Simon. 2004. Evolutionary and ecological genetics of cyclical parthenogens. Pages 122–134 in A. Moya and E. Font, eds. Evolution from molecules to ecosystems. Oxford University Press, Oxford.

Delmotte, F., N. Leterme, J. Bonhomme, C. Rispe, and J. C. Simon. 2001. Multiple routes to asexuality in an aphid species. Proceedings of the Royal Society B 268:2291–2299.

Dubey, S., J. Michaux, H. Brüinner, R. Hutterer, and P. Vogel. 2009. False phylogenies on wood mice due to cryptic cytotype-b pseudogene. Molecular Phylogenetics and Evolution 50:633–641.

Dufresne, F., and P. D. N. Hebert. 1994. Hybridization and origins of polyploidy. Proceedings of the Royal Society B 258:141–146.

Eimanífar, A., G. Van Stappen, B. Marden, and M. Wink. 2014. *Artemia* biodiversity in Asia with the focus on the phylogeography of the introduced American species *Artemia franciscana* Kellogg, 1906. Molecular Phylogenetics and Evolution 79:392–403.

Eimanífar, A., G. Van Stappen, and M. Wink. 2015. Geographical distribution and evolutionary divergence times of Asian populations of the brine shrimp *Artemia* (Crustacea, Anostraca). Zoological Journal of the Linnean Society 174:447–458.

Engelstädter, J. 2008. Constraints on the evolution of asexual reproduction. BioEssays 30:1138–1150.

———. 2017. Asexual but not clonal: evolutionary processes in automictic populations. Genetics 206:993–1009.

Estoup, A., V. Ravigne, R. Hufbauer, R. Vitalis, M. Gautier, and B. Facon. 2016. Is there a genetic paradox of biological invasion? Annual Review of Ecology, Evolution, and Systematics 47:51–72.

Glover, N. M., H. Redestig, and C. Desimone. 2016. Homoeology: what are they and how do we infer them? Trends in Plant Science. Elsevier, Oxford.

Goldschmidt, E. 1952. Fluctuation in chromosome number in *Artemia salina*. Journal of Morphology 9:111–134.

Haag, C. C. R., L. Thodosiou, J. Rjabbour-Zahab, T. Lenormand, R. Zahab, and T. Lenormand. 2017. Low recombination rates in sexual species and sex-asex transitions. Philosophical Transactions of the Royal Society B 372:20160461.

Hebert, P. D. N., and T. Crease. 1983. Clonal diversity in populations of *Daphnia pulex* reproducing by obligate parthenogenesis. Heredity 51:353–369.

Heibl, C. 2008. PHYLOCH: R language tree plotting tools and interfaces to diverse phylogenetic software packages. http://www.christophheibl.de/Rpackages.html.

Hurvich, C. M., and C. L. Tsai. 1989. Regression and time series model selection in small samples. Biometrika 76:297–307.

Jaquière, J., S. Stoeckel, C. Larose, P. Nouhaud, C. Rispe, L. Mieuzej, J. Bonhomme, et al. 2014. Genetic control of contagious asexuality in the pea aphid. PLoS Genetics 10:1–10.

Judson, O. P., and B. B. Normark. 1996. Ancient asexual scandals. Trends in Ecology and Evolution 11:41–46.

Katoh, K., K. Misawa, K. Kuma, and T. Miyata. 2005. MAFFT—a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Research 30:3059–3066.

Kearney, M. 2005. Hybridization, cladistics and geographical parthenogenesis. Trends in Ecology and Evolution 20:495–502.

Kunz, D., W. Tay, S. Elfekih, K. Gordon, and P. De Barro. 2019. Take out the rubbish: removing NUMTs and pseudogenes from the *Bemisia tabaci* cryptic species mtCOI database. bioRxiv, https://doi.org/10.1101/724765.

Lee, S. C., M. Ni, W. Li, C. Shertz, and J. Heitman. 2010. The evolution of sex: a perspective from the fungal kingdom. Microbiology and Molecular Biology Reviews 74:298–340.

Lenormand, T., J. Engelstädter, S. E. Johnston, E. Wijnker, and T. Lenormand. 2016. Evolutionary mysteries in meiosis. Philosophical Transactions of the Royal Society B 371:20160001.

Lopez, J. V., N. Yuhki, R. Masuda, W. Modi, and S. J. O’Brien. 1994. Numt, a recent transfer and tandem amplification of mitochondrial DNA to the nuclear genome of the domestic cat. Journal of Molecular Evolution 39:174–190.

Lynch, M. 1990. The similarity index and DNA fingerprinting. Molecular Biology and Evolution 7:478–484.
Origin of Asexual Brine Shrimps

Maccari, M., F. Amat, and A. Gómez. 2013a. Origin and genetic diversity of diploid parthenogenetic Artemia in Eurasia. PLoS ONE 8:e63348.

Maccari, M., F. Amat, F. Hontoria, and A. Gómez. 2014. Laboratory generation of new parthenogenetic lineages supports contagious parthenogenesis in Artemia. PeerJ 2:e439.

Maccari, M., A. Gómez, F. Hontoria, and F. Amat. 2013b. Functional rare males in diploid parthenogenetic Artemia. Journal of Evolutionary Biology 26:1934–1948.

MacDonald, G. H., and R. A. Browne. 1987. Inheritance and reproductive role of rare males in a parthenogenetic population of the brine shrimp, Artemia parthenogenetica. Genetica 75:47–53.

Maniatisi, S., A. D. Baxevanis, I. Kappas, P. Deligiannidis, A. Triantafyllidis, S. Papakostas, D. Bougiouklis, et al. 2011. Is ploidy a persevering accident or an adaptive evolutionary pattern? the case of the brine shrimp Artemia. Molecular Phylogenetics and Evolution 58:353–364.

Maynard Smith, J. 1978. The evolution of sex. Cambridge University Press, Cambridge.

Meirmans, S., P. G. Meirmans, and L. R. Kirkendall. 2012. The costs of sex: facing real-world complexities. Quarterly Review of Biology 87:19–40.

Metalli, P., and E. Ballardin. 1970. Radiobiology of Artemia: radiative effects and ploidy. Current Topics in Radiation Research Quarterly 7:181–240.

Moritz, C., W. M. Brown, L. D. Denison, J. W. Wright, D. Vyas, S. Donnellan, M. Adams, et al. 1989. Genetic diversity and the dynamics of hybrid parthenogenesis in Cnemidophorus (Teiidae) and Heteronotia (Gekkonidae). Pages 87–112 in R. M. Dawley and J. P. Bogart, eds. Evolution and ecology of unisexual vertebrates. New York State Museum, Albany, NY.

Muñoz, J., A. Gómez, A. Green, J. Figuerola, F. Amat, and C. Rico. 2010. Evolutionary origin and phylogography of the diploid obligate parthenogen Artemia parthenogenetica (Branchiopoda: Anostraca). PLoS ONE 5:e11932.

Muñoz, J., A. J. Green, J. Figuerola, F. Amat, and C. Rico. 2008. Characterization of polymorphic microsatellite markers in the brine shrimp Artemia (Branchiopoda: Anostraca). Molecular Ecology Resources 9:547–550.

Mura, G., and L. Nagorskaya. 2005. Notes on the distribution of the genus Artemia in the former USSR countries (Russia and adjacent regions). Journal of Biological Research 4:139–150.

Naganawa, H., and G. Mura. 2017. Two new cryptic species of Artemia (Branchiopoda, Anostraca) from Mongolia and the possibility of invasion and disturbance by the aquaculture industry in East Asia. Crustaceana 90:1679–1698.

Narbel-Hofstetter, M. 1964. Les alterations de la meise chez les animaux parthenogenetiques. Springer, Vienna.

Nascetti, G., P. Bondanelli, A. Aldinucci, and R. Cimmaruta. 2003. Genetic structure of bisexual and parthenogenetic populations of Artemia from Italian brackish-hypersaline waters. Oceanologica Acta 26:93–100.

Neiman, M., D. Paczesniak, D. M. Soper, A. T. Baldwin, and G. Hehman. 2011. Wide variation in ploidy level and genome size in a New Zealand freshwater snail with coexisting sexual and asexual lineages. Evolution 65:3202–3216.

Neiman, M., T. F. Sharbel, and T. Schwander. 2014. Genetic causes of transitions from sexual reproduction to asexuality in plants and animals. Journal of Evolutionary Biology 27:1346–1359.

Nougué, O., E. Flaven, R. Jabbour-Zahab, N. O. Rode, M.-P. Dubois, and T. Lenormand. 2015a. Characterization of nine new poly-morphic microsatellite markers in Artemia parthenogenetica. Biochemical Systematics and Ecology 58:59–63.

Nougué, O., N. O. Rode, R. Jabbour-Zahab, A. Ségard, L.-M. Chevin, C. R. Haag, and T. Lenormand. 2015b. Automixis in Artemia: solving a century-old controversy. Journal of Evolutionary Biology 28:2337–2348.

Otto, S. P., and T. Lenormand. 2002. Resolving the paradox of sex and recombination. Nature Review Genetics 3:252–261.

Otto, S. P., and J. Whitton. 2000. Polyploid incidence and evolution. Annual Review of Genetics 34:401–437.

Paland, S., J. K. Colbourne, and M. Lynch. 2005. Evolutionary history of contagious asexuality in Daphnia pulex. Evolution 59:800–813.

Paradis, E., and K. Schliep. 2019. ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. Bioinformatics 35:526–528.

Perey, M. L., J. R. Valverde, B. Batuecas, F. Amat, R. Marco, and R. Garesse. 1994. Speciation in the Artemia genus: mitochondrial DNA analysis of bisexual and parthenogenetic brine shrimps. Journal of Molecular Evolution 38:156–168.

Rey, O., B. Facón, J. Foucaud, A. Loiseau, and A. Estoup. 2013. Androgenesis is a maternal trait in the invasive ant Wasmannia auropunctata. Proceedings of the Royal Society B 280:20131181.

Rode, N. O., A. Charmantier, and T. Lenormand. 2011. Male-female coevolution in the wild: evidence from a time series in Artemia franciscana. Evolution 65:2881–2892.

Rode, N. O., R. Jabbour-Zahab, L. Boyer, E. Flaven, F. Hontoria, G. Van Stappen, F. Dufresne, C. Haag, and T. Lenormand. 2022. Data from: The origin of asexual brine shrimps. American Naturalist, Dryad Digital Repository, https://doi.org/10.5061/dryad.7h4j0zsb.

Sandrock, C., and C. Vorburger. 2011. Single-locus recessive inheritance of asexual reproduction in a parasitoid wasp. Current Biology 21:433–437.

Sánchez, M. I., F. Hortas, J. Figuerola, and A. J. Green. 2012. Comparing the potential for dispersal via waterbirds of a native and an invasive brine shrimp. Freshwater Biology 57:1896–1903.

Sainz-Escudero, L., E. K. López-Estrada, P. C. Rodríguez-Flores, and M. García-Paris. 2021. Setting taxonomic and nomenclatural problems in brine shrimps, Artemia (Crustacea: Branchiopoda: Anostraca), by integrating mitogenomics, marker discards and nomenclature rules. PeerJ 9:e10865.

Schliep, K. P. 2011. phangorn: phylogenetic analysis in R. Bioinformatics 27:592–593.

Schurko, A. M., M. Neiman, and J. M. Logsdon. 2009. Signs of sex: what we know and how we know it. Trends in Ecology and Evolution 24:208–217.

Shadrin, N., and E. Anufrieva. 2012. Review of the biogeography of Artemia leachi, 1819 (Crustacea: Anostraca) in Russia. International Journal of Journal of Artemia Biology 2:51–61.

Simon, P., J. Narayan, A. Houtain, A. Derzelle, L. Baudry, E. Nicolas, M. Cariou, et al. 2021. Chromosome-level genome assembly reveals homologous chromosomes and recombination in asexual rotifer Adineta vaga. Science Advances 7:eabq4216.

Simon, J., F. Delmotte, C. Rispe, and T. Crease. 2003. Phylogenetic relationships between parthenogens and their sexual relatives: the possible routes to parthenogenesis in animals. Biological Journal of the Linnean Society 79:151–163.
Swarms of diploid asexual brine shrimps (Artemia parthenogenetica) in a saltern pond (Odiel, southern Spain). The bright red color and the swarming behavior are induced by parasitic cestodes (Flamingolepis liguloides). This manipulation likely increases cestode transmission to their final host, the greater flamingo (Phoenicopterus roseus), through a greater predation of cestode-infected brine shrimps. Photo credit: Marta Sanchez.