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Abstract The parthenogenetic all-female marbled crayfish is a novel research model and potent invader of freshwater ecosystems. It is a triploid descendant of the sexually reproducing slough crayfish, Procambarus fallax, but its taxonomic status has remained unsettled. By crossbreeding experiments and parentage analysis we show here that marbled crayfish and P. fallax are reproductively separated. Both crayfish copulate readily, suggesting that the reproductive barrier is set at the cytogenetic rather than the behavioural level. Analysis of complete mitochondrial genomes of marbled crayfish from laboratory lineages and wild populations demonstrates genetic identity and indicates a single origin. Flow cytometric comparison of DNA contents of haemocytes and analysis of nuclear microsatellite loci confirm triploidy and suggest autopolyploidization as its cause. Global DNA methylation is significantly reduced in marbled crayfish implying the involvement of molecular epigenetic mechanisms in its origination. Morphologically, both crayfish are very similar but growth and fecundity are considerably larger in marbled crayfish, making it a different animal with superior fitness. These data and the high probability of a divergent future evolution of the marbled crayfish and P. fallax clusters suggest that marbled crayfish should be considered as an independent asexual species. Our findings also establish the P. fallax-marbled crayfish pair as a novel paradigm for rare chromosomal speciation by autopolyploidy and parthenogenesis in animals and for saltational evolution in general. **Key words:** marbled crayfish, autopolyploidy, parthenogenesis, epigenetics, chromosomal speciation, saltational evolution

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In the last decade, the marbled crayfish (Marmorkrebs) has gained considerable attention in the scientific community and the public because of its obligatory parthenogenetic reproduction, its suitability as a research model and its high potential as an invasive species [1-9]. It was discovered in 1995 in the German aquarium trade [2] and has become a popular pet in Europe and other continents since then [10,11]. Thriving wild populations have meanwhile developed from releases in several European countries and Madagascar and are feared to threaten native crayfish species by competition and transmission of the crayfish plague [7-9,12,13]. By comparison of morphological traits and molecular markers, Martin and colleagues [14] have identified the sexually reproducing slough crayfish *Procambarus fallax* from Florida and southernmost Georgia as the mother species of marbled crayfish. However, its taxonomic position remained unsettled. Martin et al. [14] suggested the provisional name Procambarus fallax forma virginalis, being aware that forma is not a valid category in animal taxonomy. Meanwhile, several important characteristics of marbled crayfish have been described in detail, including morphology [12], embryonic development [15,16], life history [16-19], parthenogenetic reproduction [1,20,21] and a triploid karyotype [22]. Speciation in parthenogenetic lineages is a problematic issue because parthenogens do not fit into the classical concepts of speciation, as discussed in detail by Mayr [23], Coyne and Orr [24], Barraclough et al. [25], Birky and Barraclough [26] and Martin et al. [14]. However, Barraclough and colleagues emphasized the importance of understanding diversification and speciation in asexual organisms, not least to test theories about the evolutionary advantage of sex [25,26]. They provided a theory on speciation in asexuals, which they named Evolutionary Genetic Species Concept [26]. This theory focuses on the criterion that the individuals of the parent species and the neo-species form discrete clusters of very similar genotypes and phenotypes. The new cluster should be of a single origin and both clusters

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laboratory population and the aquarium trade, (3) Procambarus alleni (Faxon, 1884) from the

2.2 Cross-breeding experiments

For the 38 crossbreeding experiments we used three *P. fallax* males with total lengths (TL=tip of rostrum to end of telson) of 3.1-5.2 cm, five *P. fallax* females with TLs of 3.5-4.2 cm, 14 marbled crayfish females with TLs of 4.0-6.3 cm and two *P. alleni* males with TLs of 5.1-5.3 cm. All males were in the reproductively competent Form I as indicated by the presence of hooks on the ischia of the 3rd and 4th peraeopods. Eight of the 14 marbled crayfish females and 4 of the 5 *P. fallax* females had well-developed glair glands on the underside of the pleon indicating ovarian maturity and receptiveness. The behavioural experiments were performed in aquaria with an area of 26x16 cm without shelter. Pairs were observed for 2 hours and copulation was regarded as successful when the partners remained in typical copulation position for more than 10 min. Parentage of the offspring was determined by microsatellite analysis.

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published mitochondrial DNA fragments from P. fallax (FJ619800) and P. alleni (HQ171462,

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internal standard. The mass transitions resulting from the loss of desoxyribose (5-

methylcytidine: 242 Th \rightarrow 126 Th, [D₃]-5-methylcytidine: 245 Th \rightarrow 129 Th) by collision induced dissociation (CID) were analysed in dynamic multiple reaction monitoring mode (DMRM). Calibration curves using a stable isotope labelled internal standard were established for quantification of 5-methylcytidine. The linear regressions resulting from the double logarithmic plots were used to correlate the respective signals from LC-MS/MS analysis to known amounts of substance. The yield of detected modification was normalized to guanosine content (as equivalent to cytidine content) because of better signal quality. To assess the amount of guanosine, the areas of the DAD results, gained during the LC analysis, were correlated to their respective amounts of substance in the same way as above.

2.7 Investigation of morphological characters and life history traits

For comparison of morphological characters between marbled crayfish and *P. fallax* we used marbled crayfish with TLs of 4.0-8.4 cm and body weights of 1.4-15.2 g and *P. fallax* females with TLs of 3.6-5.7 cm and weights of 1.1-4.5 g. We focussed on annulus ventralis (sperm receptacle), areola of the carapace, cheliped chelae and coloration, the taxonomically most relevant characters in female Cambaridae [35-37]. For comparison of life history traits we analysed growth, time of sexual maturity, body size and clutch size. Growth was determined in batches raised under the same conditions by measurement of carapace length (CL), total length (TL) and body weight. Sexual maturity was deduced from the presence of glair glands. Mean and maximum body and clutch sizes were taken from our laboratory animals and published data on wild marbled crayfish and *P. fallax*.

3. Results

3.1 Crossbreeding experiments and parentage analysis

Crossbreeding experiments were performed to investigate whether marbled crayfish and *P*.

fallax can interbreed and produce viable offspring. Behavioural observations revealed that

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products that could be used for fragment length determination in marbled crayfish and P.

fallax, namely PclG-02, PclG-04 and PclG-26. PclG-02 and PclG-26 were polymorphic and

family groups of marbled crayfish females 1-4 x *P. fallax* male 1 were identical between mothers and offspring, namely 267 bp/ 271 bp/303 bp at locus PclG-02 and 189 bp/191 bp at locus PclG-26, but differed from the allele combination of the male that was 255 bp/267 bp and 185 bp/207 bp, respectively (table 2). All measurements were repeated at least twice, and in the case of the unusual PclG-02 up to five times per specimen. Our data indicate that the male did not contribute to the genome of the offspring and that the progeny is the product of apomictic parthenogenesis. The microsatellite patterns were not only identical between mother and offspring but also between the four batches (table 2) demonstrating clonality of all marbled crayfish from our laboratory.

The *P. fallax* male 1 *x P. fallax* female 1 family was used as a positive control. Analysis of locus PclG-26 revealed the allele combinations 185 bp/207 bp in the father, 179 bp/185 bp in the mother and 179 bp/185 bp (2x), 179 bp/207 bp (4x), 185 bp/185 bp (4x) and 185 bp/207 bp (4x) in the 14 offspring. These data indicate Mendelian distribution and

demonstrate that both parents contributed equally to the genome of the offspring, as is

3.2 Single origin and clonality of marbled crayfish populations

expected for sexually reproducing species.

For a more detailed genetic analysis of marbled crayfish, we established complete mitochondrial genome sequences of specimens from our Heidelberg and Petshop lineages and from wild populations of Lake Moosweiher (Germany) and Madagascar by high-coverage shotgun sequencing and sequence mapping. Remarkably, these mitochondrial genome sequences were completely identical (figure 2), thus confirming the clonal nature of the tested populations and their single origin. Comparison of our sequences with the mitochondrial genome sequence of marbled crayfish published earlier by Shen *et al.* [44] revealed 6 scattered mismatches and major differences in one fragment ranging from position 4600 to

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5500. These differences are probably related to technical issues because Shen and colleagues used PCR-based methods and primer walking single/double strands sequencing [44] whereas we used next-generation sequencing with a sequencing coverage per nucleotide of >100x. We also established complete mitochondrial genome sequences for *P. fallax* and *P.* alleni. Analysis of the mitochondrial 12S rRNA, 16S rRNA and cytochrome oxidase subunit I genes have earlier indicated a close relationship between marbled crayfish and these species [1,7,14]. P. alleni occurs sympatrically with P. fallax in many locations in Florida [45] and was therefore regarded as a candidate that might have contributed to the origination of marbled crayfish by hybridization with *P. fallax* [46]. Sequence comparison revealed 144 single nucleotide polymorphisms (SNPs) between marbled crayfish and *P. fallax* but 1165 SNPs between marbled crayfish and P. alleni (figure 2). Interestingly, these SNPs were not evenly distributed over the mitochondrial genome, which explains why in the study by Martin et al. [14] small genetic differences between marbled crayfish and P. fallax were detected in the cytochrome oxidase subunit I gene but not in the 12S rRNA gene. Our results confirm the close genetic relationship between marbled crayfish and P. fallax and a greater distance towards P. alleni. The single origin and clonality of marbled crayfish from the laboratory and the wild was further confirmed by the analysis of microsatellite loci PclG-02, PclG-04 and PclG-26 in 24 specimens from our laboratory lineages (see parentage analysis), six specimens from a stable wild population in Lake Moosweiher [47] and one specimen from Madagascar [7]. All these marbled crayfish showed the same microsatellite patterns, namely the allele associations 267 bp/271 bp/303 bp at locus PclG-02, 159 bp at PclG-04 and 189 bp/191 bp at PclG-26. The fragment lengths of the alleles of locus PclG-02 overlapped in marbled crayfish (267-303 bp) and P. fallax (239-267 bp) but were longer in P. alleni (329-384 bp) and shorter in P. clarkii (211-228 bp). Marbled crayfish shared two of six alleles with *P. fallax*, namely 267 bp at

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locus PclG-02 and 159 bp at locus PclG-04, but none with the other species thus confirming the particularly close relationship between *P. fallax* and marbled crayfish. 3.3 Ploidy status of marbled crayfish Martin et al. [22] recently used karyological analysis to demonstrate that marbled crayfish has a triploid genome. Our microsatellite analysis confirms this finding. Marbled crayfish generally have the allele association 267 bp/271 bp/303 bp at locus PclG-02 (figure 3a), whereas P. fallax, P. alleni and P. clarkii have one or two alleles at this locus, which is consistent with diploid and sexually reproducing species. In an earlier paper, Martin et al. [20] have also analysed locus PclG-02 and reported only two alleles of 267 bp and 271 bp. However, a recent re-examination of their material confirmed the presence of the third 303 bp allele (G. Scholtz, personal communication). We further corroborated triploidy in marbled crayfish by flow cytometric measurement of the DNA content of haemocytes in marbled crayfish and P. fallax. Haemocytes are particularly suitable for this purpose because they are devoid of somatic polyploidization [48]. Our results showed a significant 1.4-fold higher DNA content in the blood cells of marbled crayfish (figure 3b), which is consistent with triploidy. 3.4 Comparison of DNA methylation between marbled crayfish and *Procambarus fallax* In order to test if the marbled crayfish and P. fallax clusters also differ with respect to epigenetic markers we determined global DNA methylation by mass spectrometry in identically raised and age and size-matched representatives of both crayfish. DNA methylation represents a widely conserved epigenetic mark that is often associated with polyphenism and adaptive phenotypic changes [49,50]. Comparison of three juveniles and selected organs (hepatopancreas, abdominal musculature and ovary) of three adults revealed a consistently and highly significantly reduced level of DNA methylation in marbled crayfish

when compared to *P. fallax* (figure 4). The ten *P. fallax* samples together had a DNA methylation level of 2.93±0.15% (mean ± standard deviation) whereas the ten marbled crayfish samples together had a level of only 2.40±0.08%. These results suggest that marbled crayfish have a considerably different DNA methylation pattern.

3.5 Comparison of morphological characters between marbled crayfish and *P. fallax*

Comparison of the most relevant taxonomic characters of cambarid females [35-37] between marbled crayfish and *P. fallax* corroborated the high degree of morphological similarity between the two crayfish as previously established by Kawai *et al.* [12] and Martin *et al.* [14]. The diagnostically most meaningful trait in females of the genus *Procambarus* is the annulus ventralis, which is bell-shaped with a tilted S-shaped sinus in both marbled crayfish and *P. fallax* (figure 5a,b). This typical form is not found in other *Procambarus* species [37] as best exemplified by the differently shaped sperm receptacle of the closely related *P. alleni* (figure 5c). The areola, an unpaired structure on the dorsal midline of the carapace, is also very similar in marbled crayfish and *P. fallax* with respect to shape and length-to-width proportion (figure 5d,e). The same holds for the cheliped chelae, which closely resemble each other in both crayfish in shape, dentation and setation (figure 5f,g), and the coloration pattern, which consists of distinct marmorated spots and dark dorsolateral stripes on carapace and pleon (figure 5h,i). Size, form and coloration of the marmoration spots are highly variable not only in the sexually reproducing *P. fallax* but also in the genetically uniform marbled crayfish as a result of stochastic developmental variation [21,51].

3.6 Comparison of life history traits between marbled crayfish and *P. fallax*

In contrast to the morphological characters, life history features like growth and fecundity are markedly different between marbled crayfish and *P. fallax*. Figure 6 gives an example for differences in the speed of growth between identically raised laboratory populations of the

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same age. At day 250 after hatching, when the first females in both crayfish had reached sexual maturity, mean body weight was almost twice as large in marbled crayfish as in P. fallax females. Maximum body and clutch sizes were also markedly higher in marbled crayfish. The largest specimen in our laboratory had a carapace length of 4.9 cm, a total length of 10.3 cm and a body weight of 30.1 g (figure 7a). In the wild, the largest of the 1084 marbled crayfish measured [7,12,47, M. Pfeiffer and C. Chucholl, personal communication] was found in Lake Moosweiher and had a CL of 4.9 cm and a weight of 32.0 g [47]. In contrast, the largest of the 4710 wild P. fallax examined [36,52-54] had a CL of only 3.4 cm, corresponding to a TL of 7.4 cm and a weight of approximately 11.5 g. The largest clutches of marbled crayfish in the laboratory and the wild consisted of 731 eggs (figure 7b) and 724 eggs [47], respectively, which is 5.6 fold higher than the largest clutch of 130 eggs reported for *P. fallax* in literature [53]. The analysis of life history features of the slough crayfish by van der Heiden [54] corroborated that *P. fallax* reaches only rarely a size of more than 6.5 cm TL. The differences in growth and fecundity between marbled crayfish and *P. fallax* were also confirmed by the analysis of published data for egg-carrying females from comparable climatic regions. Ovigerous marbled crayfish from Madagascar had a mean CL of 3.5 cm, a mean TL of 7.4 cm and a mean clutch size of 300 eggs [7], whereas ovigerous P. fallax from the Everglades National Park in Florida had a mean CL of 1.8 cm, a mean TL of 3.8 cm and a mean clutch size of 41 eggs only [53], indicating that body size and fecundity is significantly increased in marbled crayfish (figure 7c,d). These findings identify important phenotypic differences between marbled crayfish and *P. fallax* that have not been recognized previously. 4. Discussion Our results demonstrate that marbled crayfish meets all criteria for asexual speciation [25-28].

It is separated from the mother species, *P. fallax*, by reproductive isolation, significant

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genomic and epigenetic differences and superior life history traits. Our data further support a single origin. In addition, all populations known to date live outside the natural range of P. fallax, suggesting geographical isolation. They are unified in one cluster by common phenotypic, genetic and epigenetic characteristics, despite their broad geographical distribution. These commonalities and differences towards P. fallax make it very likely that the marbled crayfish and slough crayfish clusters will evolve differently, which is the main criterion for erecting an asexual species [26]. Martin et al. [14] have previously suggested that marbled crayfish should be considered as an independent species when a single origin and/or regional populations in the wild have been established. Our findings clarify the former issue and provide additional evidence for cytogenetic, genetic and phenotypic differences between marbled crayfish and *P. fallax*. As such, marbled crayfish should now be named *Procambarus* virginalis, as suggested previously [14]. The formal description of marbled crayfish as a new species will be detailed in a separate publication. Marbled crayfish appeared first in 1995 in the German aquarium trade. Thereafter, aquarists have propagated it in captivity, and since about 2003, releases have resulted in the establishment of thriving wild populations in Central Europe and Madagascar [5,7-9,12,47]. The "mega-population" [46] in innumerable aquarium tanks on various continents and the known wild populations are apparently all descendants of the single clone or single individual that was introduced in Germany in 1995. Our results confirm this single origin by the identity of the mitochondrial genomes and microsatellite patterns in samples of captive and wild populations. One of the samples analysed in our study, the Heidelberg specimen, can be directly traced back to the year 1995 and to the oldest marbled crayfish for which written records exist (F. Steuerwald, personal communication). It is unknown whether marbled crayfish emerged in the natural range of P. fallax or in captivity. Scholtz [4], Faulkes [5] and Martin [46] summarized possible scenarios for the first alternative including hybridization with coexisting *Procambarus* species and geographic

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parthenogenesis. These authors and Chucholl [9] also stressed that in captivity there were many more candidates for hybridization than the naturally coexisting six *Procambarus* species [36,52] because crayfish were popular pets already in the 1990s. Faulkes [5] emphasized that all surveys on P. fallax in Florida and Southern Georgia revealed males and females arguing against the presence of pure marbled crayfish populations in the natural range of P. fallax. Moreover, none of the articles on wild P. fallax [36,45,52-54] mentioned specimens above 7.4 cm TL, which would again support the absence of primary populations of marbled crayfish. In sympatric populations, small and medium-sized marbled crayfish and P. fallax females would be indistinguishable by morphological criteria alone. However, by the use of genetic markers marbled crayfish could now be identified. Particularly useful is the highly specific tri-allelic microsatellite locus PclG-02, which could be assayed in large samples with reasonable expenditure. However, time for the detection of primary populations may be limited because marbled crayfish are already widespread in American aquaria [11] and their release into the natural range of *P. fallax* would render the search for primary populations of marbled crayfish impossible. Our crossbreeding experiments with marbled crayfish, P. fallax and P. alleni revealed that marbled crayfish and *P. fallax* still recognize each other as sexual partners but not marbled crayfish and P. alleni. Recognition of sexual partners in crayfish is mainly based on chemical signatures of the urine but may also include visual and tactile cues [38,39]. Marbled crayfish and P. fallax copulate readily with each other. However, the progeny of such pairings are pure marbled crayfish resulting from parthenogenesis. These findings demonstrate reproductive isolation and suggest that the reproductive barrier is set at the cytogenetic rather than the behavioural level. Mechanical barriers can be largely excluded because the sperm receptacles are structurally very similar in marbled crayfish and *P. fallax* females and because we have repeatedly observed insertion of the male gonopods into the annulus ventralis of marbled crayfish. We attempted to directly prove sperm transfer by analysing moulted sperm

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receptacles of females that had successfully produced offspring. However, we did not find any sperm remnants neither in marbled crayfish nor *P. fallax* females. The morphological features and microsatellite patterns strongly suggest that marbled crayfish originated by autopolyploidization and not by hybridization with a closely related species, which is by far the most frequent cause of triploidy in animals [55-58]. Typically, hybrids between two crayfish species are clearly recognizable because of their intermediate morphological characters [59,60]. However, marbled crayfish do not show such hybrid features [12,14, this study]. Conversely, autopolyploids are usually morphologically similar to their diploid progenitors [61], and the morphological similarity between marbled crayfish and P. fallax is therefore consistent with autopolyploidization. There is also no evidence for hybridization on the genetic level and no strong bias towards heterozygosity in the microsatellite pattern, which would be typical for hybrids [62,63]. Of the seven microsatellite loci that were investigated in marbled crayfish so far, three were homozygous and four were heterozygous [20,21, this study], thus largely excluding allopolyploidization for marbled crayfish. Furthermore, Martin and colleagues have recently shown that the nuclear elongation factor 2 (EF-2) gene is identical in marbled crayfish and P. fallax but differs from other Procambarus species like P. alleni, P. clarkii, P. acutus and P. liberorum [22]. These findings provide additional support for the origin of marbled crayfish by autopolyploidization. We admit that the presence of three alleles, as observed in locus PclG-02 in marbled crayfish, can be interpreted to reflect an origin by hybridization. However, such a pattern can also occur in autopolyploids, namely when an unreduced diploid egg is fertilized by a sperm from the same species, or alternatively, by simultaneous fertilization of a haploid egg by two sperms with different alleles. In shrimp, fish and bivalve aquaculture, autopolyploid triploids with tri-allelic loci are artificially produced by the prevention of polar body I extrusion in

fertilized eggs either by temperature shock or chemicals like 6-dimethylaminopurine [64,65].

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Marbled crayfish may thus have arisen by a heat or cold shock in the sensitive phase of egg development in a captive *P. fallax* female, possibly during transportation. The origin of parthenogenesis in marbled crayfish is probably a by-product of polyploidization but the causal relationship of polyploidy and parthenogenesis is not yet understood [46]. Infectious parthenogenesis by the feminizing bacterium Wolbachia, which is widespread in crustaceans [66], was excluded by the use of molecular probes for the parasite [2]. In plants, it was shown that polyploidy per se can have an immediate impact on the reproductive biology of a species [67]. In animals, however, obligate parthenogenesis is relatively rare. It has been described in some asexual invertebrate families and a few vertebrate hybrids [26,68-71] and is mostly associated with allopolyploidy. Autopolyploidy is much less common and is usually not associated with parthenogenesis, perhaps with the exception of some high arctic ostracods and polyploid populations of the brine shrimp Artemia parthenogenetica [72,73]. Artificially induced autopolyploid shrimp and fish are usually sterile [74], making the combination of autopolyploidy and parthenogenesis in marbled crayfish rather unique. Polyploids often have life history traits that are different from those of the parent species. Growth, number of offspring and other quantitative traits can either be decreased or increased when compared to the diploid ancestors [75-77]. In marbled crayfish, growth, maximum body size and fecundity were significantly increased when compared to P. fallax, whereas the time of sexual maturity was similar (7,36,47,54, this study). Longevity may also be increased in marbled crayfish. Maximum age so far recorded is 1610 days in marbled crayfish [19] and 980 days in *P. fallax* (Z. Faulkes, personal communication). These superior fitness traits, together with parthenogenetic reproduction, are probably causative for the remarkable success of marbled crayfish as an invasive species in Central Europe and Madagascar [7-9,47]. Chucholl [9] calculated an almost double FI-ISK (Freshwater Invertebrate Invasiveness Scoring Kit) score for marbled crayfish when compared to *P. fallax*, making it a high risk

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species for Central Europe. Moreover, Feria and Faulkes [78] predicted with climate and habitat based Species Distribution Models that marbled crayfish could inhabit a larger geographical area than its mother species P. fallax when released in the southern states of the USA, thus illustrating the ecological superiority of marbled crayfish. In allopolyploids, the increase of life history traits is usually explained as the result of heterozygosity, which is well known as heterosis effect or hybrid vigor [79,80]. However, this explanation is not applicable for autopolyploids because autopolyploidization enhances only the copy number of already existing genes. However, novel traits do not necessarily require new genes or new developmental pathways to come into being but can instead arise from recruitment of already existing developmental processes into new contexts [81,82]. Thus, trait alteration in marbled crayfish may have been caused by altered gene dosage, the rearrangement of gene-networks and the modulation of gene expression by changes in epigenetic regulation. Changes in epigenetic regulation can be deduced from the significantly reduced level of global DNA methylation in marbled crayfish when compared to P. fallax. DNA methylation is an epigenetic mechanism that considerably affects plant and animal phenotypes [49,50,83]. It is responsive to environmental and genomic stresses including polyploidization [50] and might thus contribute to speciation in polyploids. In plants, the increase or reduction of global DNA methylation after autopolyploidization is well known [61,84]. It is also well established that DNA methylation and other epigenetic mechanisms contribute to the establishment of reproductive barriers [85,86] and the expression of hybrid vigor in allopolyploid plants [87]. In marbled crayfish, epigenetic mechanisms may thus have been involved in the acquisition of novel fitness traits. Chen et al. [88] reported that polyploidization is often accompanied or followed by intense rearrangements in the genome, which stabilize the new lineage. These rearrangements, which are associated with epigenetic changes, can include loss of DNA. For example, in synthetic

autopolyploids of annual phlox, *Phlox drummondii*, an immediate loss of 17% of total DNA has been observed with a further reduction of up to 25% upon the third generation [89]. Such mechanisms may also have operated during transition from *P. fallax* to marbled crayfish and might explain why triploid marbled crayfish have only a 1.4-fold rather than a 1.5-fold increased DNA content when compared with its diploid mother species.

Speciation by autopolyploidization is a special case of chromosomal speciation that is well-known in plants [61] but virtually unknown in animals. Chromosomal speciation is a complementary concept to the better known speciation by changes in allele frequency distribution and can result in the almost instantaneous production of new species and phenotypic novelty within one generation [90-92]. This "saltational speciation" or "saltational evolution" [93-95] has largely been ignored by gradualism-based Modern Synthesis, which may be due to its rarity in animals, the lack of mechanistic understanding and the dearth of suitable models. Marbled crayfish represents a contemporary animal example of autopolyploid speciation, which likely started about 20-30 generations ago. Comparative genome and epigenome sequencing approaches will be required to fully understand the genetic and epigenetic differences between both species.

5. Conclusion

Marbled crayfish can be regarded as a new species that originated from *P. fallax* by triploidization and concomitant epigenetic alterations, as shown by our combined morphological, behavioural, genetic and epigenetic analysis. Marbled crayfish is morphologically very similar to its mother species but has superior fitness traits. Genetic data suggest an instantaneous speciation by autopolyploidization and parallel change of the mode of reproduction from gonochorism to parthenogenesis. The young evolutionary age of marbled crayfish, which is possibly three decades or less, may offer the possibility to identify key events for this type of speciation. The combination of autopolyploidy and obligate

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parthenogenesis is common in plants but very rare in animals. Thus, the *P. fallax*-marbled crayfish pair provides an interesting new model system to study asexual speciation and saltational evolution in animals and to determine how much genetic and epigenetic change is necessary to create a new species. **Acknowledgement.** We thank Michael Pfeiffer (Gobio, March-Hugstetten, Germany) and Christoph Chucholl (Fisheries Research Station Baden-Württemberg, Langenargen, Germany) for providing marbled crayfish from Lake Moosweiher and for information on the biology of marbled crayfish in this lake, Frank Glaw (Zoologische Staatssammlung, Munich, Germany) and Miguel Vences (Braunschweig University of Technology, Germany) for the Madagascar sample, the Bundesamt für Umwelt (Bern, Switzerland) for the *Procambarus* clarkii samples, Frank Steuerwald (KABS, Waldsee, Germany) for information on the oldest known marbled crayfish, Chris Lukhaup (Hinterweidenthal, Germany) for figure 5i, Thomas Carell (Ludwig-Maximilians-University, Munich, Germany) for providing [D₃]-dm⁵C internal standard for mass spectrometry, Günter Raddatz and Carine Legrand (DKFZ) for statistical help, the DKFZ Flow Cytometry and Genomics and Proteomics Core Facilities for flow cytometry and DNA sequencing services, and Gerhard Scholtz (Humboldt University, Berlin, Germany), Bronwyn W. Williams (North Carolina Museum of Natural Sciences, Raleigh, USA) and Zen Faulkes (University of Texas-Pan American, Edinburg, USA) for valuable comments that improved the manuscript. **Authors' contributions.** G.V. conceived of the study, participated in the design of the study, sampled the tissues, performed the cross-breeding experiments and analysed the morphological and life history data; C.F. carried out the assembly and analysis of mitochondrial genome sequences and the determination of DNA contents by flow cytometry; K.H. maintained laboratory crayfish cultures and prepared DNA samples; A.S., J.P. and R.S.

- performed the analysis of the microsatellite markers; K.S and M.H. carried out the mass
- spectrometric measurement of DNA methylation; F.L. participated in the design of the study
- and coordinated the study. G.V. and F.L. wrote the manuscript. All authors revised the
- manuscript and gave final approval for publication.
- Data accessibility: The mitochondrial DNA sequences have been deposited in GenBank
- under the accession numbers KT074363, KT074364 and KT074365.
- **Ethics statement:** All crayfish experiments were performed by approval of the institutional
- animal welfare committee, in compliance with local standards and guidelines.
- 548 **Competing interests:** We have no competing interests.
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Males	Marbled crayfish females										P. fe	P. fallax females							
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	P1	P2	Р3	P4	P5
P. fallax 1	х	х	х	х	XX		х	xo	o	х		х		o	х	х			X
P. fallax 2						o	х		o	х	o	х					o	х	
P. fallax 3								х					x			х			х
P. alleni 1					О		О	00											0
P. alleni 2							00	00											0

x: mating; o: no mating; two letters: results of two trials.

Table 2. Parentage analysis in crossbreeds of marbled crayfish *x P. fallax*.

Specimens	Microsatellite loci					
	PclG-02	PclG-26				
P. fallax father 1	255/267	185/207				
Marbled crayfish mothers 1-4	267/271/303	189/191				
Offspring of mother 1 (n=6)	267/271/303	189/191				
Offspring of mother 2 (n=5)	267/271/303	189/191				
Offspring of mother 3 (n=6)	267/271/303	189/191				
Offspring of mother 4 (n=3)	267/271/303	189/191				

Values indicate fragment lengths in base pairs.

Figures and figure legends



Figure 1. Mating of marbled crayfish female with *P. fallax* male. The male (top) holds the female firmly with the chelipeds and ischial hooks and his gonopods are plugged into the female's spermatheca.

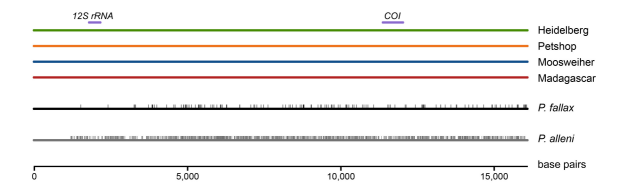


Figure 2. Comparison of complete mitochondrial genomes of marbled crayfish, *P. fallax* and *P. alleni*. The sequences of marbled crayfish from two laboratory populations (Heidelberg, Petshop) and two wild populations (Moosweiher, Madagascar) are completely identical. In contrast, the sequences of *P. fallax* and *P. alleni* differ in 144 and 1165 SNPs (vertical lines) from marbled crayfish, respectively. Purple bars indicate positions of *12S rRNA* and *cyto-chrome oxidase subunit I* (COI) genes that were earlier used for phylogenetic analysis [14].

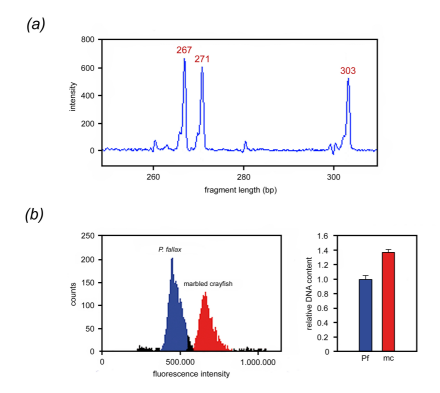


Figure 3. Ploidy status of the marbled crayfish genome. (a) Microsatellite locus PclG-02 in marbled crayfish showing a combination of three alleles of 267 bp, 271 bp and 303 bp fragment length. (b) Flow cytometry of haemocytes of P. fallax (Pf) and marbled crayfish (mc) revealing an approximately 1.4 fold increased DNA content in marbled crayfish. The right panel shows the means and standard deviations of two biological and three technical replicates. Differences are highly significant (p=1.33x10⁻⁷, Welsh two-sided t-test).

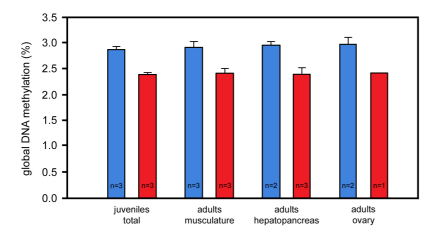


Figure 4. Differences in global DNA methylation between marbled crayfish (red) and P. fallax (blue). Analysed were three complete juveniles and major organs of three adult females in each crayfish. Note consistently and significantly greater methylation levels in P. fallax (p=1.48x10⁻⁷ for the sum of all samples, Welsh two-sided t-test). Error bars: standard deviations.

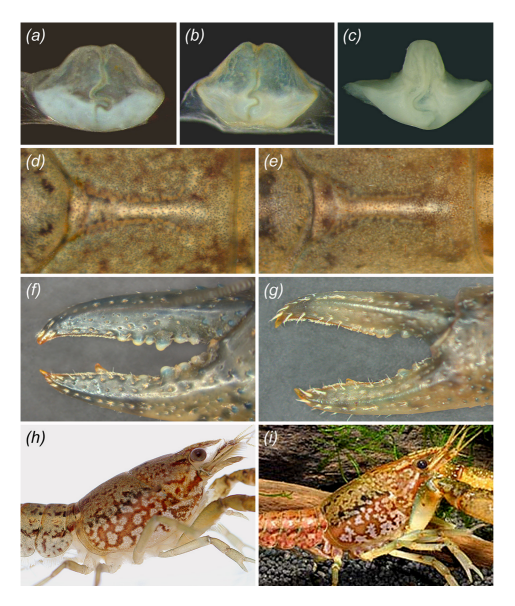


Figure 5. Comparison of morphological characters between marbled crayfish and *P. fallax*. (a) Annulus ventralis from exuvia of marbled crayfish. (b) Annulus ventralis of *P. fallax*. (c) Annulus ventralis of *P. alleni*. Note striking structural difference to sperm receptacles of marbled crayfish and *P. fallax*. (d) Areola of marbled crayfish. (e) Areola of *P. fallax*. (f) Left cheliped of marbled crayfish of 8.4 cm TL. (g) Left cheliped of *P. fallax* female of 4.7 cm TL. Form, dentation and setation of the chelae are very similar in both species. (h) Coloration of cephalothorax in marbled crayfish. (i) Coloration of cephalothorax in *P. fallax* male (photo: C. Lukhaup).

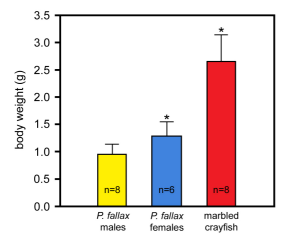


Figure 6. Comparison of growth between marbled crayfish and *P. fallax*. The three groups were reared for 250 days at 20°C under identical conditions and fed with the same food *ad libitum*. The differences between marbled crayfish and *P. fallax* females are highly significant (asterisks; $p=2.06\times10^{-5}$; Welsh two-sided t-test). Error bars: standard deviations.

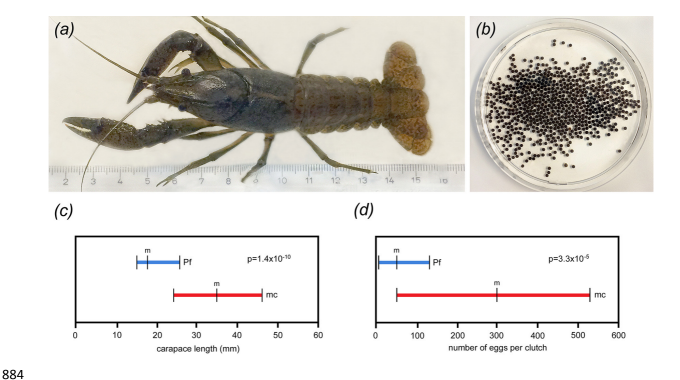


Figure 7. Comparison of body size and fecundity between marbled crayfish and *P. fallax.* (a) Largest marbled crayfish from our laboratory having a total length of 10.3 cm. (b) Clutch of same specimen consisting of 731 eggs. (c) Differences in carapace length between populations of ovigerous marbled crayfish (mc) and *P. fallax* females (PF) from comparable climatic regions. Data for marbled crayfish (n=57) was obtained in Madagascar [7] and data for *P. fallax* (n=27) was obtained in Florida [53]. Horizontal bars indicate ranges and vertical lines indicate mean values (m) and lower and upper range limits. The difference between marbled crayfish and *P. fallax* females is highly significant as indicated by the p-value. (d) Differences in clutch size between the same populations as in (c). The difference is highly significant as indicated by the p-value. For statistical calculations, the standard deviation was taken as half the range, and a Bonferroni adjustment for multiplicity was applied.