Chromosomal location of rDNA clusters and TTAGG telomeric repeats in eight species of the spittlebug genus *Philaenus* (Hemiptera: Auchenorrhyncha: Aphrophoridae)

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**Abstract.** A cytogenetic investigation was performed in eight species of the spittlebug genus *Philaenus* using silver-NOR (AgNOR)-banding and fluorescence in situ hybridization (FISH) with 18S rDNA and (TTAGG), telomeric probes. This is the first application of FISH technique in the Auchenorrhyncha, a suborder of the Hemiptera. FISH along with the rDNA probe revealed differences between species in the number and chromosomal location of major ribosomal RNA gene sites, the so-called nucleolar organizer regions (NORs). However, we found a lack of perfect correlation between the results of AgNOR-staining and rDNA-FISH in the detection of NORs. FISH with the telomeric probe confirmed that the chromosome ends of the *Philaenus* species are composed of the (TTAGG)ₙ nucleotide sequence, which is a common motif of insect telomeres.

**INTRODUCTION**

The chromosomes of Auchenorrhyncha have been studied cytologically for over 100 years (see Halkka, 1959), although they are not particularly amenable to cytological investigation, being holokinetic and hard to differentiate in a karyotype. Some progress toward the identification of separate marker chromosomes in the karyotypes of holokinetic groups was achieved using differential staining techniques (different types of chromosome banding); however, only a few publications of this kind have addressed Auchenorrhyncha specifically to date (Perepelov et al., 2002; Kuznetsova et al., 2003, 2009, 2010; Maryańska-Nadachowska et al., 2008, 2012). During the past few decades, fluorescence in situ hybridization (FISH) has become an important method in insect comparative cytogenetics. This technique allows genes or DNA sequences to be directly located on chromosomes in cytological preparations, although as yet, no such data are available for Auchenorrhyncha.

In Auchenorrhyncha, there are a few genera in which the chromosome number has been subjected to change during the course of speciation, a prime example being the spittlebug genus *Philaenus* (Maryańska-Nadachowska et al., 2012). Presently, a total of eight species are recognized in this genus, including the Mediterranean species, *P. tesselatus*, *P. loukasi*, *P. arslani*, *P. signatus*, *P. maghresignus*, *P. tarifa*, and *P. italosignus*, and the Holarctic species *P. spumarius* (Drosopoulous & Remane, 2000). While the Mediterranean species are sympatric with *P. spumarius*, they are partially allopatric with one another. Mainly due to the outstanding dorsal colour polymorphism, the genus *Philaenus* has long received the widespread attention of biologists. Among other aspects, several studies have dealt with chromosome and molecular analyses of this group (Kuznetsova et al., 2003; Maryańska-Nadachowska et al., 2008, 2010, 2011, 2012; Seabra et al., 2010). Molecular analysis has confirmed the monophyly of the genus and the species status of all species with the exception of *P. tesselatus*, which has been suggested as representing a geographical and morphological form of *P. spumarius* (Maryańska-Nadachowska et al., 2010, 2011). Cytogenetic analysis using standard staining, C-banding, AgNOR-banding, and fluorochrome banding (CMA₃ and DAPI) revealed that the species differ from one another in terms of chromosome number, sex chromosome system and certain other characters. Altogether, three different chromosome numbers (2n = 20, 23, and 24 for males) and three variations of the male sex chromosome system (X0, XY, and X₁X₂Y) have been established in the genus *Philaenus* (Maryańska-Nadachowska et al., 2012).

The present study represents part of an ongoing project exploring the genus *Philaenus*. Here, we applied FISH with an 18S rDNA probe to investigate eight *Philaenus* species. Our aim was to identify additional chromosome markers for a comparative study of representatives of the genus. Similarly, we examined the presence of TTAGG telomeric repeats in *Philaenus* species using FISH. So far, the only study to have reported the presence of (TTAGG)ₙ sequence in the genome of *Callipygona pellucida* (Delphacidae) is that of Frydrychová et al. (2004). Our study is thus the first cytogenetic investigation of Auchenorrhyncha using the FISH technique.
TABLE 1. Species, collection localities, number of males and host plants of the eight *Philaenus* species studied.

<table>
<thead>
<tr>
<th>Species</th>
<th>Data and locality of collection</th>
<th>No. of males examined</th>
<th>Host plants</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. spumarius</em></td>
<td>July 2007, Bieszczady Mts, SE Poland</td>
<td>7</td>
<td>polyphagous</td>
</tr>
<tr>
<td><em>P. tesselatus</em></td>
<td>June 2005; S Portugal</td>
<td>7</td>
<td>polyphagous</td>
</tr>
<tr>
<td><em>P. loukasi</em></td>
<td>August 2004 and July 2005; mountains in Greece</td>
<td>8</td>
<td><em>Eryngium</em> sp.</td>
</tr>
<tr>
<td><em>P. arslani</em></td>
<td>June–July 2005 Mazraat Kfar Zebrani (Kesrouane), Lebanon</td>
<td>5</td>
<td><em>Eryngium</em> sp.</td>
</tr>
<tr>
<td><em>P. italosignus</em></td>
<td>May 2002 and 2006; Sicily, Italy</td>
<td>12</td>
<td><em>Asphodelus aestivus</em></td>
</tr>
<tr>
<td><em>P. maghresigus</em></td>
<td>June 2005; S Spain</td>
<td>6</td>
<td><em>A. aestivus</em></td>
</tr>
<tr>
<td><em>P. signatus</em></td>
<td>August 2005; July 2008; central Greece</td>
<td>8</td>
<td><em>A. aestivus</em></td>
</tr>
<tr>
<td><em>P. tarifa</em></td>
<td>June 2005; S Spain</td>
<td>6</td>
<td><em>A. aestivus</em></td>
</tr>
</tbody>
</table>

MATERIAL AND METHODS

Eight *Philaenus* Stål, 1864 species were used in the present study. Specimens of seven species (*P. tesselatus* Melichar, 1889; *P. loukasi* Drosopoulos & Asche, 1991; *P. arslani* Abdoul-Nour & Lahoud, 1996; *P. signatus* Melichar, 1896; *P. maghresigus* Drosopoulos & Remane, 2000; *P. tarifa* Remane & Drosopoulos, 2001; and *P. italosignus* Drosopoulos & Remane, 2000) were collected during the 2002–2007 field seasons in the Mediterranean region, while specimens of *P. spumarius* (Linnaeus, 1758) originated from Poland. The species collection sites and host plants as well as the number of specimens studied are given in Table 1. On capture, insects were immediately fixed in Carnoy fixative (3 parts of 96% alcohol, 1 part of glacial acetic acid) and stored at −15°C until required. Preparations were made from testes, which were dissected in a drop of 45% acetic acid and squashed under a coverslip on a glass microscope slide. The slides were frozen using dry ice, the coverslips were removed with a razor blade, and the preparations then air dried.

Ag-staining

Silver staining was performed following the technique of Howell & Black (1980). Slides were treated in 50% silver nitrate (AgNO₃) with gelatin as the developer in a moist chamber at 65°C for 8 min. After the preparations had been stained appropriately, the reaction was stopped by rinsing three times with water, and the preparations air dried.

Fluorescence in situ hybridization (FISH) with 18S rDNA and (TTAGG)ₙ telomeric probes

For rDNA-FISH, a 1.8 kb fragment of 18S rDNA was generated by PCR using genomic DNA isolated from the specimens of *Phyrrocoris apterus* as a template. Amplification was performed in a final volume of 25 µl containing 1.5 mM MgCl₂, 2.5 mM dNTPs, 10 µM of both the 18Sai forward primer (5'-CCT GAG AAA CGG CTA CCA CAT C-3') and the 18Sbi reverse primer (5'-GAG TCT CGT TCG TTA TCG GA-3') (Whiting et al., 1997), 100 ng template DNA, and 5 U Taq DNA polymerase (Qiagen, Hilden, Germany). An initial denaturation period of 3 min at 94°C was followed by 30 cycles of 60 s at 94°C, annealing for 60 s at 51°C, a 1.5 min extension step at 72°C, and concluded with a final extension step of 10 min at 72°C. The PCR product was then labelled with digoxigenin-11-dUTP during additional PCR cycles.

For FISH with both rDNA and telomeric probes, we used the procedure described in Lichter et al. (1988). Chromosome preparations were treated with 100 µg/ml RNase A. After denaturation, the chromosomes were hybridized with a probe mixture containing 20 ng of 18S rDNA probe, 20–50 ng telomeric probe (TTAGG), and 10 µg of sonicated salmon sperm DNA (Invitrogen, Life Technologies Inc., San Diego, CA, USA). Hybridization signals were detected with avidin-Alexa 488 (Invitrogen) and mouse anti-digoxigenin antibodies conjugated to Cy3-IgG (Jackson Immuno Research Laboratories, West Grove, PA, USA). The preparations were counterstained with 2 µl/ml DAPI (4'-6'-diamidino-2-phenolindole) (Sigma-Aldrich, St. Louis, MO, USA) for 15 min and then mounted in an antifade based on DABCO (1,4-diazabicyclo[2.2.2]octane; Sigma-Aldrich) (see Sahara et al., 1999). At least 10 meiotic divisions (diplotene/ diakinesis) per male and 5 to 12 males per species were analyzed using the FISH and AgNO₃ techniques (Table 1). Photographs were taken with a Nikon Eclipse 400 light microscope fitted with a CCD DS-U1 camera using Chroma filter sets (for FISH). Lucia Image 5.0 software was applied and images were processed in Adobe Photoshop Elements 9.

RESULTS

Standard karyotypes

In agreement with our previous results (Maryańska-Nadachowska et al., 2012), the eight studied *Philaenus* species showed karyotypes with a male meiotic formula of n = 11 + X₀ in *P. spumarius* and *P. tesselatus*, n = 9 + neo-XY in *P. loukasi* and *P. arslani*, n = 11 + neo-XY in *P. signatus*, *P. maghresigus*, and *P. tarifa*, and n = 10 + neo-neo-X,X,Y in *P. italosignus* (Figs 1–22).

Detection of a tandem telomeric repeat sequence by FISH with a (TTAGG)ₙ probe

In each of the species studied, bright fluorescent signals were seen at the ends of spermatocyte chromosomes (Figs 2, 5, 7, 9, 12, 14, 15, 18, 21, 22). These findings indicate conclusively that the telomeres in *Philaenus* species are composed of the (TTAGG)ₙ nucleotide sequence.

Location and activity of ribosomal genes revealed by FISH with an 18S rDNA probe and silver staining

A difference in ribosomal gene locations was found between *P. spumarius* and *P. tesselatus*, which share a common chromosomal complement (2n = 22 + X). In *P.
*spumarius*, the silver staining of male meiotic cells labelled NORs in the largest pair of autosomes (no. 1) as well as in the medium-sized pair (presumed no. 6) (Fig. 1). The FISH experiments confirmed the presence of rDNA arrays in the same bivalents (Fig. 2). In *P. tesselatus*, silver staining predominantly detected Ag-positive sites in the autosome pair no. 1. However, an additional Ag-positive site was occasionally observed in a medium-sized bivalent (presumed pair no. 6) in some cells (Figs 3 and 4). In contrast to silver staining, FISH experiments performed in this species invariably showed 18S rDNA only in the largest pair of autosomes (Fig. 5). In both species, rDNA clusters (i.e. NORs) were located terminally. Hybridization signals were uniform in size and intensity with the exception of the large NOR-bivalent in *P. spumarius*, which was constantly heteromorphic for the size of the signals (Fig. 2).

In yet another pair of species sharing the same gross structure of karyotype, *P. loukasi* and *P. arslani* (2n = 18 + neo-XY), a difference between ribosomal gene locations was also found. In *P. loukasi*, silver staining revealed two Ag-positive regions in male meiotic cells (Fig. 6). Unfortunately, exact identification of the NOR-bearing chromosomes was not possible with silver staining. rDNA-FISH clearly detected a bright signal on the Y chromosome in the neo-XY bivalent and a weak signal on one of the medium-sized pairs of autosomes (presumed no. 6) (Fig. 7). The signals were seen close to the ends of the NOR-chromosomes. In *P. arslani*, silver staining likewise revealed two Ag-positive regions in male meiotic cells, one of them being of uncertain location, the other one clearly belonging to the largest pair of autosomes (Fig. 8). However, using rDNA-FISH the only signal was observed on the largest autosome bivalent (Fig. 9). The

Figs 1–9. Chromosomal locations of Ag-positive sites (asterisks; Figs 1, 3, 4, 6, 8), clusters of 18S rDNA (green signals; Figs 2, 5, 7, 9) and TTAGG telomeric repeats (red signals; Figs 2, 5, 7, 9) in males of *Philaenus* species. 1 and 2: *P. spumarius*. 1 – diplotene with Ag-positive sites in the largest bivalent and in a medium-sized bivalent; 2 – diakinesis with two rDNA signals in the same two bivalents (signals in the largest bivalent are heteromorphic in size). 3–5: *P. tesselatus*. 3 – pachytene with Ag-positive sites in the largest bivalent and in one of the smaller bivalents; 4 – diplotene with the only Ag-positive site in the largest bivalent; 5 – diakinesis with two rDNA signals in opposite ends of the largest bivalent. 6 and 7: *P. loukasi*. 6 – pachytene with Ag-positive sites, one in the largest bivalent and the other of uncertain location; 7 – diakinesis with rDNA signals in the Y chromosome of the neo-XY bivalent and in a medium-sized bivalent (arrowhead). 8 and 9: *P. arslani*. 8 – pachytene with two Ag-positive sites, one (interstitial) in the largest bivalent and the other of uncertain location; 9 – diplotene with the only interstitial rDNA signal in the largest bivalent. Bar = 10 µm.
The special feature of this karyotype is that the rDNA array is most probably subterminal, being universally seen at a short distance from the end of the NOR-chromosome, as evidenced by both silver-staining and FISH (Figs 8, 9).

In *P. signatus*, *P. maghresignus*, and *P. tarifa*, another three species sharing a common chromosomal complement (*2n = 22 + neo-XY*), an essential difference in ribosomal gene location was found. In *P. signatus*, silver
staining revealed one (Fig. 10), or sometimes two (Fig. 11), Ag-positive sites in prophase I meiotic cells. One of the sites was always located in a medium-sized bivalent, whereas the location of the other site, when present, was not reliably identified. rDNA-FISH invariably detected a single rDNA array in one of the medium-sized bivalents, probably no. 6 (Fig. 12). In *P. maghresignus*, two silver-positive sites were seen in pachytene cells (Fig. 13). rDNA-FISH detected rDNA arrays in the sex chromosome bivalent, in which one signal was terminal in the neo-Y chromosome, while the other was interstitial in the neo-X chromosome (Figs 14, 15). In *P. tarifa*, silver staining revealed a single pair of NOR-chromosomes in mitotic prophases (Fig. 16); however, two autosome bivalents showed AG-positive sites in meiotic cells (Fig. 17). rDNA-FISH revealed two rDNA clusters in this karyotype, one located in the larger (probably the largest) bivalent, the other one in a medium-sized bivalent, probably no. 6 (Fig. 18).

In *P. italosignus* (2n = 20 + X1Y1X2Y2), two chromosomes with Ag-positive sites were found in mitotic prophases (Fig. 19). In meiotic prophases, one or sometimes two,
Ag-positive sites were observed (Fig. 20). The 18S rDNA probe localized two clusters in the sex chromosome trivalent in meiotic cells; one of these signals was terminal in the neo-X₂, whereas the other was interstitial in the neo-Y chromosome (Figs 21, 22).

It is worth noting that the Ag-positive sites could be seen and identified only in the extended prophase chromosomes. At mitotic and meiotic prophase, argentophilic material was separated into interconnected granules grouped more commonly around particular (presumably NOR-bearing) chromosomes.

**DISCUSSION**

In the genus *Philaenus*, differences in chromosome number (Maryańska-Nadachowska et al., 2012; present paper) are largely in agreement with the recognized grouping proposed on the basis of morphology (Drosopoulou & Remane, 2000) as well as being supported by larval food plant relationships (Drosopoulou, 2003). The species feeding on common asphodel, *Asphodelus aestivus*, that is, *P. signatus*, *P. maghreensis*, and *P. tarifa*, sharing 22 + neo-XY, and *P. italosignus*, having 2n = 20 + neo-neo-X₁X₂Y, are all similar in morphology and have the same or closely related karyotypes (that of *P. italosignus* has evidently arisen from 22 + neo-XY via an additional fusion between the ancestral Y chromosome and an autosome pair). These species fall into the species group “*signatus*”, defined on the basis of the morphology of male genitalia (Drosopoulou & Remane, 2000). The species *P. loukasi* (2n = 18 + neo-XY) and *P. arslani* (2n = 18 + neo-XY), whose larvae develop on host plants growing in arid conditions, share similar karyotypes, as do the polyphagous species *P. spumarius* (2n = 22 + X) and *P. tessellatus* (2n = 22 + X), feeding on a wide range of dicotyledonous plants. In turn, all of these *Philaenus* species fall into another morphologically substantiated group, “*spumarius*”. In phytophagous insects, host plant shifting might be an important factor in sympatric divergence and speciation (Berlocher & Feder, 2002; Horn et al., 2006; Loxdale, 2010). In some aphids, another group of the order Hemiptera, polymorphism for chromosome number was found to be correlated with change of host plants (Brown & Blackman, 1988; De Barro et al., 1995; Hales et al., 2006). Such karyotypic divergence in relation to host plant is known to occur in the genus *Amphorophora* where *A. idaei* Börner, which feeds on raspberry (*Rubus idaeus*) has 2n = 18, whereas *A. rubi* (Kaltenbach), which feeds on blackberry (*Rubus fruticosus agg.*) has 2n = 20. These host-specific taxa are distinguishable morphologically using multivariate analysis and presently recognized as separate species (Blackman et al., 2000). Another example concerns the closely related forms of the corn-leaf aphid, *Rhopalosiphum maidis* (Fitch). In this species there is a strong association between chromosome number and host plant. Specimens with 2n = 8 are found on *Sorghum* and *Zea mays*, whereas on northern hemisphere barley, *Hordeum vulgare*, 2n = 10 is common (Brown & Blackman, 1988).

With this assumption, we speculated that changes in the host-plant associations of *Philaenus* species may well be accompanied by chromosome rearrangements and that karyotype changes may have occurred independently several times in the evolution of the genus. Moreover, we inferred ways of chromosome rearrangement that probably occurred during the karyotype evolution of this group (Maryańska-Nadachowska et al., 2012). Assumptions of the sequence of karyotype rearrangements were, however, based on a limited set of characters, such as chromosome number, sex chromosome system, and chromosome size. Our present study was aimed to find additional markers in *Philaenus* (location of rDNA and telomeric DNA sequences revealed by FISH) for testing the validity of the aforementioned assumptions.

Indeed, 18S rDNA-FISH enabled us to detect differences in the number and location of ribosomal loci between the species, and more importantly, differences were found between species with the same chromosome number. Within the genus, three species (*P. tessellatus*, *P. arslani*, *P. signatus*) were shown to have one cluster of rDNA genes, whereas five species (*P. spumarius*, *P. tarifa*, *P. italosignus*, *P. loukasi*, *P. maghreensis*) had two clusters. In *P. italosignus* and *P. maghreensis*, rDNA sites are borne by sex chromosomes, whereas in the other species examined, by autosomes or (in *P. loukasi*) by both autosomes and the neo-Y chromosome (Fig. 23).

It is clear that the major rDNA distribution patterns show no correlation with the aforementioned grouping of the genus. *Philaenus* species evidently originate (monophyletically) from a common ancestor, as it has been confirmed by molecular analyses (Maryańska-Nadachowska et al., 2010, 2011). Differences as presently revealed suggest that karyotype changes may have occurred during species formation and evolved independently. At least four independent translocation events occurred in the evolution of the neo-XY system in *Philaenus* species. In *P. loukasi*, *P. arslani*, *P. maghreensis*, *P. signatus*, and *P. tarifa*, neo-sex chromosome systems possibly originated by fusion of a pair of autosomes with the original X chromosome, while in *P. italosignus*, the original neo-Y chromosome probably fused with the homologue of another autosomal pair, resulting in a neo-neo-X₂,Y (Maryańska-Nadachowska et al., 2012). Our present data suggest that the fused autosomes were different in every case, lacking NRs, as in *P. signatus* and *P. tarifa*, while retaining NRs in other species.

The differences observed between the very closely related *P. spumarius* and *P. tessellatus* are also noteworthy. Our molecular analyses using nucleotide sequences from two mitochondrial genes (*COI* and *Cyt B*) and one nuclear region (ITS 2) strongly support the species status of all species, the only exception being *P. tessellatus*, which may represent a geographical and morphological form of *P. spumarius* (Maryańska-Nadachowska et al., 2010, 2011). Even so, we found that these species were indeed different in that *P. tessellatus* had a single rDNA cluster in the largest autosome pair, whereas in *P. spu-
marius, the 18S rDNA probe also mapped to the largest pair of autosomes and, in addition, to one of the medium-sized bivalents.

It is of interest that autosomally located rDNA sites, when present in the species studied, were situated similarly on the largest and/or on a medium-sized bivalent. Likewise, in P. loukasi one of the two loci was located on a medium-sized bivalent. By convention, the medium-sized NOR-bivalent, when present in a species, could be identified as belonging to chromosome pair no. 6. Thus, it appears that the NOR-bearing medium-sized autosome pairs are homologues in different species. Be that as it may, the currently available chromosome markers are insufficient to test the validity of this hypothesis.

NOR-chromosomes in the species studied mostly showed terminally located rDNA clusters, but in some cases the clusters were clearly positioned in the interstitial position: in the neo-X of P. maghresignus, the neo-Y of P. italosignus, or in the autosomal NOR-bivalent of P. arsleni. These interstitial patterns argue that some ancestral NOR-bearing autosomes were involved in fusions during Philaenus phylogeny. In most species and specimens examined to date, hybridization signals were uniform in size and intensity; although two exceptions were observed. One of these exceptions was P. spumarius, in which the larger NOR-bivalent appears to be heteromorphic due to the size of the signals, while the other was P. loukasi, in which the signals on the Y chromosome and on the medium-sized pair of autosomes differed in intensity. Most probably the observed differences reflect variation in the copy number of rRNA genes either between different chromosome pairs or between two homologous chromosomes. This can be attributed to various mechanisms, including unequal crossing over, transposition, tandem amplification and other rearrangements involving homologous segments, yielding structural modification of NORs (Castro et al., 2001) The heteromorphism in size and intensity of NORs has been frequently reported in aphids (Monti et al., 2011 and reference therein), heteropterans (Grozeva et al., 2010), orthopterans (Warchałowska-Śliwa et al., 2009), as well as certain other insect orders.

It is generally accepted that NORs are often highlighted by fluorescent staining with chromomycin A3 (CMA3) that binds to GC-rich regions. In Philaenus species, we found that the position of the major rDNA sites corresponded in most cases to active Ag-NOR sites. Nevertheless, there were some rDNA-FISH clusters that were negative for Ag-NORs and CMA3 staining and rDNA sites has been reported in many animal species (Gromicho et al., 2005 and references therein), including insects such as ants (Lorite et al., 1997), beetles (Colomba et al., 2000) and grasshoppers (Vitturi et al., 2008). Thus, our data on Philaenus species support the view that not only NORs, but also some other regions, are stained with CMA3 and silver nitrate, and that some rDNA sites are not always detected using these techniques (Gromicho et al., 2005). Although our study represents the first application of FISH to Auchenorrhyncha chromosomes, it testifies that this approach is a potentially valuable tool for the physical mapping of major rDNA regions on the chromosomes of such insects.

We examined the presence of TTAGG telomeric repeats by using FISH in all species of Philaenus available to us. To date, a single attempt was made to detect telomeric repeats in Auchenorrhyncha chromosomes, this concerning the leafhopper, Calligrypona pellucida (F.) (Delphacidae) (Frydrychová et al., 2004). This particular earlier study was, however, done using Southern hybridization of genomic DNA, a technique known to rank below FISH in terms of accuracy of telomeric repeat detection, since it reveals only the presence of the sequence in the genome, but not its location. Our FISH experiments with the (TTAGG)n probe have shown that this sequence occurs at the ends of all chromosomes, providing conclusive evidence that in Philaenus species, telomeres consist of TTAGG tandem repeats. This telomere motif is present in the vast majority of evolutionary lineages of insects and other arthropods and is thought to represent an ancestral sequence of telomeres in the Arthropoda as a whole (Sahara et al., 1999; Frydrychová et al., 2004; Vitková et al., 2005; Lukhtanov & Kuznetsova, 2010). Our data on eight species of the genus Philaenus suggest that the (TTAGG)n telomeric repeat motif has been conserved in the Auchenorrhyncha.

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