A standard photomap of the ovarian nurse cell chromosomes for the dominant malaria vector in Europe and Middle East *Anopheles sacharovi*

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**Abstract**

**Background:** *Anopheles sacharovi* is a dominant malaria vector species in South Europe and the Middle East which has a highly plastic behaviour at both adult and larval stages. Such plasticity has prevented this species from eradication by several anti-vector campaigns. The development of new genome-based strategies for vector control will benefit from genome sequencing and physical chromosome mapping of this mosquito. Although a cytogenetic photomap for chromosomes from salivary glands of *An. sacharovi* has been developed, no cytogenetic map suitable for physical genome mapping is available.

**Methods:** Mosquitoes for this study were collected at adult stage in animal shelters in Armenia. Polytene chromosome preparations were prepared from ovarian nurse cells. Fluorescent in situ hybridization (FISH) was performed using PCR amplified probes.

**Results:** This study constructed a high-quality standard photomap for polytene chromosomes from ovarian nurse cells of *An. sacharovi*. Following the previous nomenclature, chromosomes were sub-divided into 39 numbered and 119 lettered sub-divisions. Chromosomal landmarks for the chromosome recognition were described. Using FISH, 4 PCR-amplified genic probes were mapped to the chromosomes. The positions of the probes demonstrated gene order reshuffling between *An. sacharovi* and *Anopheles atroparvus* which has not been seen cytologically. In addition, this study described specific chromosomal landmarks that can be used for the cytotaxonomic diagnostics of *An. sacharovi* based on the banding pattern of its polytene chromosomes.

**Conclusions:** This study constructed a high-quality standard photomap for ovarian nurse cell chromosomes of *An. sacharovi* and validated its utility for physical genome mapping. Based on the map, cytotaxonomic features for identification of *An. sacharovi* have been described. The cytogenetic map constructed in this study will assist in creating a chromosome-based genome assembly for this mosquito and in developing cytotaxonomic tools for identification of other species from the Maculipennis group.

**Keywords:** *Anopheles sacharovi*, Mosquito, Cytogenetic map, Fluorescence in situ hybridization

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Background

Anopheles sacharovi, Favr [1] is one of the Palearctic members of the Maculipennis group of malaria mosquitoes that also includes Anopheles atroparvus, Anopheles artemievi, Anopheles beklemishevi, Anopheles daceae, Anopheles labranchiae, Anopheles maculipennis, Anopheles martinius, Anopheles melanoon, Anopheles messeae, and Anopheles persiensis [2]. Four of these 11 species, An. atroparvus, An. labranchiae, An. messeae, and An. sacharovi, are considered dominant malaria vectors in Eurasia [3]. Among them, An. sacharovi has the most southern distribution and is the most dangerous malaria vector that transmits malaria in South Europe and in the Middle East [3]. Currently, An. sacharovi is involved in transmission of vivax malaria in Iran [4–6], Iraq [7] and Turkey [8]. After the collapse of the Soviet Union, this species became responsible for malaria re-emergence in Georgia [9], Armenia [10, 11] and Azerbaijan [12]. The success of An. sacharovi as a malaria vector is a result of a highly plastic adaptation of this species at both adult and larval stages [3]. It can breed in different water reservoirs, such as swamps, marshes, river margins, streams, pools, and ditches. Female mosquitoes have opportunistic blood-feeding behaviour and can feed on any available host including human, cow, sheep, chicken, horse, and donkey. It has been demonstrated that An. sacharovi is resistant to DDT [13] and dieldrin [5]. Such ecological and behavioural plasticity, together with the emerged insecticide resistance, prevented elimination of An. sacharovi during several anti-vector campaigns conducted in Israel, Greece and Turkey. Additionally, global warming raises concerns about the possible spread of An. sacharovi to the dry areas and territories with high altitudes [14].

An importance of An. sacharovi as a malaria vector stimulates studies of this species from different perspectives including cytogenetic analyses. The first cytogenetic map for this species was developed based on photo images of lacto-aceto-orcein-stained polytene chromosomes from salivary glands [15]. This map was used for species identification of An. sacharovi and for the cytotaxonomic validation of the species status of An. martinius Shingarev originally described in Uzbekistan in 1926 [16]. Anopheles martinius is identical to An. sacharovi based on external characters including egg morphology, and its name for a long period of time was considered a synonym for An. sacharovi [17]. The cytogenetic analysis clearly demonstrated that An. sacharovi and An. martinius differ from each other by two fixed paracentric inversions on chromosomes X and 3L and, thus, represent two different species [15]. Moreover, polytene chromosomes of laboratory hybrids between An. sacharovi and An. martinius demonstrated partial asynapsis of the homeologous chromosomes, which is typical for the interspecies hybrids in other species of Diptera [18]. The importance of this finding was emphasized by G. B. White in his review of systematic reappraisal of the An. maculipennis complex [19]. Later, the species validity of both An. sacharovi and An. martinius was supported by fixed nucleotide substitutions in the internal transcribed spacer 2 (ITS2) [20]. Chromosome polymorphism has not been yet detected in natural populations of An. sacharovi.

In addition to species diagnostics, the cytogenetic map of An. sacharovi was utilized to study chromosome evolution and phylogeny of Palearctic members of the Maculipennis group [21]. The analysis of overlapping chromosomal inversions in 7 sibling species from the An. maculipennis complex revealed three branches of the phylogenetic tree: An. atroparvus-An. labranchiae; An. melanoon-An. maculipennis-An. messeae; and An. sacharovi-An. martinius. Among them, the branch An. atroparvus-An. labranchiae was considered as the basal. Although the phylogeny based on ITS2 sequences supported the presence of 3 major clades in the Palearctic group, the molecular analysis suggested that An. sacharovi is the basal lineage [20, 22, 23]. Thus, the phylogeny of the Maculipennis group remains unresolved.

The genomics era offers new opportunities for the development of modern genome-based strategies for vector control [24, 25] including the CRISPR/Cas9-based gene-drive technologies [26]. Following the major malaria vector in Africa Anopheles gambiae [27], the genomes for other malaria mosquitoes have been sequenced [28–30]. The availability of cytogenetic maps allows the development of high-quality genome assemblies anchored to the chromosomes. However, only 5 chromosome-based genome assemblies have been developed for malaria mosquitoes [28, 29, 31, 32], including one for the dominant vector of malaria in Europe An. atroparvus [33]. The comparison of the chromosome-based assemblies provided important insights into chromosomal evolution in the genus Anopheles. A high rate of the sex chromosome evolution [34], whole-arm translocations among autosomes [35], and inter-arm rearrangements [32] have been reported.

In this study, a standard cytogenetic photomap for An. sacharovi was developed. The suitability of this map for physical mapping was demonstrated by placing five PCR-amplified genes to the chromosomes of this species. In addition, cytogenetic landmarks that can be used for cytotaxonomic identification of An. sacharovi were described. This new map will assist in the development of the chromosome-based genome assembly for this important malaria vector.
Methods
Mosquito collection
About 250 adult specimens of *An. sacharovi* were collected in September 2015 in the horse and pig shelters in Arakavan village, Ararat region, Armenia (39°59’58.5″N, 44°27’38.9″E). Ovaries from half-gravid females were dissected, placed in individual 0.5-ml tubes with cold Carnoy’s fixative solution (ethanol: acetic acid in 3:1 ratio), and stored at −20 °C.

Chromosome preparation
Ovarian nurse cells polytene chromosome preparation was done following a standard procedure [33, 36]. A piece of ovary was placed on the slide into a drop of 50% propionic acid for 10 min, covered by the coverslip and squashed. Chromosome preparations with a high-quality banding pattern were selected for the image acquisition and fluorescence in situ hybridization (FISH). High-quality preparations were dipped in liquid nitrogen, the coverslips were removed, and the preparations were dehydrated in an ethanol series (50, 70 and 100%), air dried and stored at room temperature.

Chromosome map development
Images for the chromosome map were obtained using a phase contrast AxioImager A1 microscope (Carl Zeiss, OPTEC LLC, Novosibirsk, Russia), CCD camera MrC5, and AxioVision 4.8.1 software (Carl Zeiss, OPTEC LLC, Novosibirsk, Russia) at the 100x magnification. The images were combined, straightened, shaped, and cropped using Adobe Photoshop CS2 [33, 36]. The marking of nurse cell chromosomes by divisions and sub-divisions was conducted using a previously developed salivary glands chromosome map [15].

PCR of DNA probes
Mosquito DNA was extracted from specimens using standard protocol for Qiagen DNeasy Mini Kit (Qigenc, Germantown, MD, USA). Gene-specific primers were designed to amplify four unique exon sequences using PRIMER-BLAST software available at the NCBI website [37]. Genes from different chromosome arms of *An. sacharovi* were selected based on similarity with the *An. atroparvus* genome [28, 33, 36]. The primer design was based on gene annotations from the AatrE1 genome assembly available at the VectorBase website [38]. PCR was performed in the presence of 1xPCR buffer (SibEnzyme Ltd., Novosibirsk, Russia), 2.5 mM magnesium chloride (SibEnzyme Ltd., Novosibirsk, Russia), 0.2 mM dNTP (Thermo Fisher Scientific TM, USA), and 0.02 u/μl Taq Polymerase (SibEnzyme Ltd., Novosibirsk, Russia).

DNA-probe labelling and fluorescence in situ hybridization (FISH)
Gene-specific DNA-probes were labelled using a Random Primer Labeling Protocol: 25 µl of labelling reaction contained 50-ng DNA, 1x Klenow buffer (Thermo Fisher Scientific TM, USA), 44-ng/μl Exo-Resistant Random Primer (Thermo Fisher Scientific TM, USA) 0.1-mM dATP, dGTP, and dCTP and 0.015-mM dTTP, 0.016-mM TAMRA-5-dUTP, and 5 units of Klenow fragment (Thermo Fisher Scientific TM, USA) in a PCR tube. The required amounts of DNA, Klenow buffer, and Random Primers were mixed, brought up to 12 µl with water, and heated at 95 °C for 5 min in a thermocycler. The solution was chilled on ice, and appropriate amounts of nucleotides, Klenow fragment and water were added to reach 25 µl. The reaction mix was incubated at 37 °C for 18 h. FISH was performed using previously described standard protocol [39, 40].

Results
The development of a standard cytogenetic map for *Anopheles sacharovi*
In this study, a high-quality photomap was developed for the polytene chromosome from the ovarian nurse cells of a dominant malaria vector in Europe and Middle East, *An. sacharovi*. As was shown before [15], chromosome complement of *An. sacharovi* consists of 3 pairs of chromosomes in diploid cells that correspond to 5 polytene chromosome arms in ovarian nurse cells (Fig. 1). The X chromosome is represented by a single arm. Arms of chromosome 2 are usually connected together, whereas arms of chromosome 3 get separated from each other during squashing of chromosomal preparations because of fragile connections between them [41, 42]. Relative lengths of the chromosomes in *An. sacharovi* (Table 1) are similar to that in *An. atroparvus* [33]. The X chromosome is the shortest arm, and the 3R arm is the longest arm in the polytene chromosome complement. The banding pattern and quality of *An. sacharovi* chromosomes are similar to that in *An. atroparvus* [33].

To develop a cytogenetic map for *An. sacharovi*, ~150 chromosome preparations from ovarian nurse cells of 76 specimens were screened for the presence of high-quality polytene chromosomes with clear banding structure. Ten to 15 images of each chromosome arm were selected and further processed for map development. Chromosome images were combined, straightened, shaped, and cropped using Adobe Photoshop CS2 [33, 36]. Following the nomenclature previously developed for the salivary gland chromosomes of *An. sacharovi* [15], ovarian nurse
Chromosomes were sub-divided into 39 numeric divisions and 119 lettered sub-divisions (Fig. 2). The order of lettered sub-divisions for the 2L and 3L arms in the new map was changed because in the previous map the order of lettered sub-divisions was opposite to the order of numbered divisions. A set of three dark narrow bands in region 10A and 2 dark bands in region 10B can be considered an additional landmark for the middle of 2R arm chromosome. The telomere region of the 2L arm has a long light area limited by 2 thin bands in region 21C and a dark band in region 21B.

In contrast to the light-diffused pericentromeric region in 2R arm, 2L arm contains two dark bands in region 15A–15B.

In contrast to the telomere ends in chromosome 2, the telomere end in chromosome arm 3R is striped, and it starts with a dark band in region 22A followed by 6 additional distinct bands in region 22A–23B. Pericentromeric region 32B–32C of 3R arm is asynaptic in ~60% nuclei, but homologous chromosomes are always paired in region 32B that contains a dark band. Three dark bands in regions 26A, 26B and 26C that localize at an equal distance from each other and are followed by a thin, light region in 27BC and can also be used for recognizing the 3R arm. Chromosome arm 3L is the shortest autosomal arm that, in contrast to 3R arm, has a flared light telomere end. The pericentromeric region usually forms a fan consisting of diffuse chromatin fibrils. In addition, 3L contains a typical landmark for all members of the Maculipennis group in region 38A–38B, known as 'bird eye', represented by a set consisting of a dark crescent-shaped band followed by a dot-like band.

Physical mapping of orthologous genes to Anopheles sacharovi chromosomes

To test the utility of the new chromosome map developed for An. sacharovi, four probes designed based on An. atroparvus genes were hybridized to the ovarian nurse cell chromosomes of An. sacharovi using FISH. The genes AATE009010, AATE009656, AATE006419, and AATE016383 were selected from the scaffolds previously mapped to the X chromosome, 2R arm, 3R arm, and 3L arm of An. atroparvus, respectively [32, 33]. Each DNA probe hybridized to the chromosomes of An. sacharovi (Fig. 3), and their positions were placed on the chromosome map (Fig. 2). Gene AATE009010 was mapped to region 1B in the chromosome X, gene AATE009656 was assigned to region 7B in the 2R arm, gene AATE006419 was anchored to region 23C/24A in the 3R arm, and gene...
AATE016383 was placed in the region 35B in the 3L arm. Thus, all markers were successfully mapped to a specific position in the newly developed map of *An. sacharovi* chromosomes, confirming the utility of this map for physical mapping.

**A cytotoxicnomic approach to identification of *Anopheles sacharovi***

Polytene chromosomes can serve as cytotoxicnomic tools for species identification [21]. The cytogenetic map for *An. sacharovi* is the third map among the recently developed maps for members of the Maculipennis group. The other two maps were developed for *An. atroparvus* [33] and *An. beklemishevi* [43]. The comparison of polytene chromosomes among the species allowed to identify species-specific features that can be used for cytotoxicnomic diagnostics. The overall quality of the banding pattern in the polytene chromosomes of the ovarian nurse cells was species-specific (Fig. 4). For example, chromosomes of *An. sacharovi* and *An. atroparvus* are characterized by fuzzy boundaries between bands and interbands. In contrast, chromosomes of *An. beklemishevi* have a clear banding pattern with sharp band-interband boundaries that are similar to the structure of the salivary gland chromosomes. Although all 3 species have similar landmarks in the middle of the arms, the positions and orientations of these landmarks in the chromosomes vary among species because of the presence of fixed pericentric inversions. For interspecies comparison, traditional nomenclature was followed in which the banding pattern of *An. atroparvus* is considered as standard for the entire species group [21]. The most robust landmark for the species diagnostics is a ‘bird eye’ landmark in the 3L arm. This landmark is organized as a tandem of two dark bands: a dot-like band and a dark crescent-shaped band (Fig. 4d). In *An. atroparvus*, the ‘bird eye’ is located close to the centromere in a standard orientation: the dot-like band is followed by the crescent-shaped band. By contrast, in *An. sacharovi*,
the 'bird eye' is located closer to the telomere and has an inverted orientation. In *An. beklemishevi*, this landmark moved almost to the middle of the arm and has a standard orientation. Figure 4d shows the positions of the inversion that differentiates *An. sacharovi* and *An. martinius*. In the latter species, the 'bird eye' is located even closer to the telomere and has a standard orientation. The landmarks on chromosomes 2R (Fig. 4b) and 3R (Fig. 4c) have reverse orientations in *An. beklemishevi* in comparison with both *An. atroparvus* and *An. sacharovi*. The re-arrangements in the X chromosome (Fig. 4a) and 3L arm (Fig. 4d) between the species are seen based on the position of 2 genes, AATE009010 and AATE016383, that are located on opposite sides of the chromosomes in *An. sacharovi* and *An. atroparvus*.

In addition to fixed chromosomal inversions between the species, pericentromeric regions of the chromosomes have distinct species-specific features in *An. sacharovi*, *An. atroparvus* and *An. beklemishevi* [41]. For example, both *An. sacharovi* and *An. beklemishevi* have a compact dot-like pericentromeric band in the X chromosome, whereas in *An. atroparvus*, the X chromosome has three thin line-shape bands (Fig. 5A). On the other hand, the pericentric regions of the X chromosome in *An. atroparvus* and *An. sacharovi* consist of a diffuse chromatin, whereas this part of the X chromosome in *An. beklemishevi* has a regular banding pattern. The homologous chromosomes 2 in *An. sacharovi* and *An. atroparvus* are paired in pericentromeric regions. The pericentric region in 2R and 2L arms in *An. beklemishevi* is asynaptic, and
the arms of chromosome 2 have a fragile connection because of their attachment to the nuclear envelope [42] that is usually disrupted by squashing during chromosome preparation (Fig. 5B). Similarly, fragile pericentric regions are characteristics of chromosome 3 in all 3 species. However, in *An. beklemishevi*, this region does not form a fan-like structure as is seen in *An. sacharovi* and *An. atroparvus* (Fig. 5C). Thus, this study demonstrated that cytogenetic maps could serve for the cytotaxonomic diagnostic of *An. sacharovi* and other species from the Maculipennis group.

**Discussion**

This study developed a high-resolution map for unstained polytene chromosomes from the ovarian nurse cells of *An. sacharovi*. Chromosome images were developed using phase-contrast microscopy and high-resolution digital photography that provide a detailed banding pattern of the polytene chromosomes. The first chromosome map of *An. sacharovi* was developed for the polytene chromosomes from the chromosomes of salivary glands [15]. Those chromosomes were stained with lacto-aceto-orcein and imaged using film photography. That technique provided too contrasted banding patterns of the chromosomes because of the overstaining of the large bands under staining of thin bands. Also, using ovarian nurse cell chromosomes in this study helped to avoid the loss of detail in the structure of the pericentromeric regions, usually affected by under-replication in salivary glands due to their heterochromatic nature [44].

The cytogenetic map that was developed for *An. sacharovi* is one of 12 chromosome maps constructed for anopheline species using digital imaging technology, since 2000 (Table 2). A special emphasis has been placed on cytogenetic map construction for the dominant vectors of malaria from different parts of the world [3, 45].
Cytogenetic maps are now available for the malaria vectors in Africa: *An. gambiae* [36], *Anopheles funestus* [46] and *Anopheles nili* [35]; in Asia: *An. stephensi* [47] and *Anopheles sinensis* [48]; and in America: *Anopheles albimanus* [49] and *Anopheles darlingi* [50, 51]. Chromosome maps have been used to identify and describe inversion polymorphism in natural populations of *An. funestus* [46], *An. nili* [35] and *An. darlingi* [50, 51]. Physical maps of microsatellite markers have been used in population genetic studies of *An. funestus* [52], *An. stephensi* [53] and *An. nili* [54]. Although inversion polymorphism has not been reported for *An. sacharovi*, the availability of the cytogenetic map may help to discover polymorphic inversions in this species in the future. Considering the limited number of population studies conducted for *An. sacharovi*, as well as the large geographical range of this mosquito, spreading from southern Europe to the Middle East, such discovery is quite possible.

Another reason that stimulated the recent interest in cytogenetic research in mosquitoes is the need to develop chromosome-based physical genome maps to enhance the quality of genome assemblies (Table 2). A superior physical genome map has been developed based on polytene chromosomes for the African

### Table 2: The development of chromosome maps for malaria mosquitoes

| Mosquito species | Chromosome source | Cytogenetic map year/reference | Physical map year/reference | Genome map year/reference |
|------------------|-------------------|--------------------------------|-----------------------------|---------------------------|
| An. albimanus     | Salivary glands   | 2000 [49], 2017 [32]           | 2000 [49]                   | 2017 [32]                |
| An. atroparvus    | Ovarian nurse cells | 2015 [33]                      | 2009 [57]                   | 2015 [33], 2018 [55]     |
| An. beklemishevi  | Ovarian nurse cells | 2018 [43]                      |                              |                          |
| An. darlingi      | Salivary glands   | 2010 [50], 2016 [51]           |                              |                          |
| An. funestus      | Ovarian nurse cells | 2001 [46]                      | 2004 [52]                   | 2002 [58]                |
| An. gambiae       | Ovarian nurse cells | 2010 [36]                      |                              |                          |
| An. lesteri       | Salivary glands   | 2016 [59]                      | 2016 [59]                   |                          |
| An. nili          | Ovarian nurse cells | 2011 [35]                      | 2011 [35], 2012 [54]       |                          |
| An. owengensis    | Ovarian nurse cells | 2013 [60]                      |                              |                          |
| An. sacharovi     | Ovarian nurse cells | Current study                  |                              |                          |
| An. sinensis      | Salivary glands   | 2014 [48]                      | 2014 [48]                   | 2017 [56]                |
| An. stephensi     | Ovarian nurse cells | 2006 [47]                      | 2010 [39], 2010 [34], 2011 [53] | 2014 [29]  |
malaria vector An. gambiae [36]. The genome of this species has been mapped by in situ hybridization of ~2000 bacterial artificial chromosome (BAC) clones [27]. The map has been improved by additional mapping of cDNA probes that allowed a better coverage of the heterochromatic regions [31]. Finally, the genome coordinates have been placed on the cytogenetic photomap developed using a high-pressure technique for squashing chromosomal preparation [36]. This map assigns 84.3% of the AgamP3 assembly to the chromosomes with coordinates spaced at 0.5–1 Mb intervals. A remarkably high-coverage genome map has been developed for the Neotropical vector of malaria An. albimanus [32] and for the European vector An. atroparvus [55]. These maps include 98.2 and 89.6% of their genome assemblies, respectively, and represent the most complete genome maps developed for mosquitoes to date. Lower coverage physical maps have been developed for dominant Asian malaria vectors, An. stephensi [29, 39] and An. sinensis [56], and for a major malaria vector in Africa, An. funestus [28]. In this study, physical mapping was performed for 4 orthologous genes on An. sacharovi chromosomes, for which location in the An. atroparvus genome and chromosomes is known. This study demonstrates that the An. sacharovi map is suitable for physical mapping and can serve as a tool for the development of a high-quality genome assembly for this species.

In addition, chromosome banding patterns of the An. sacharovi map were compared with other chromosome maps that were recently developed for the members of the Maculipennis group, An. atroparvus [33] and An. beklemishevi [43]. The comparison clearly demonstrated that the chromosome map of An. sacharovi can be utilized as a tool for the cytotoxic features for identification of An. sacharovi and 2 other species from the Maculipennis group: An. atroparvus and An. beklemishevi, were identified and described. The cytogenetic map constructed in this study will help to create a chromosome-based genome assembly for this mosquito and will further stimulate the development of genomic-based strategies for vector control.

### Conclusions

This study reports the development of a standard photomap for the ovarian nurse cell chromosomes of An. sacharovi, one of the most dangerous malaria vectors in Europe and Middle East. The suitability of this map for physical mapping is demonstrated by successful positioning of 4 DNA probes on the map using FISH. In addition, cytotoxic features for identification of An. sacharovi and 2 other species from the Maculipennis group: An. atroparvus and An. beklemishevi, were identified and described. The cytogenetic map constructed in this study will help to create a chromosome-based genome assembly for this mosquito and will further stimulate the development of genomic-based strategies for vector control.

### Abbreviations

- DNA: deoxyribonucleic acid; dATP: deoxyadenosine triphosphate; dGTP: deoxyguanosine triphosphate; dCTP: deoxycytidine triphosphate; dTTP: deoxythymidine triphosphate; FISH: fluorescence in situ hybridization; ITS2: internal transcribe spacer 2; PCR: polymerase chain reaction.

### Authors’ contributions

IVS and MVS conceived and designed the experiments. GA, AIV, and SB performed the experiments. GA, SB, GHK, SAA, and MSA collected mosquitoes. VNS provided resources and made contributions to conception and discussion of the study. GA, IVS, and MVS conducted data analysis and wrote the manuscript. All authors read and approved the final manuscript.

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### Competing interests

The authors declare that they have no competing interests.

### Availability of data and materials

Not applicable.

### Consent of publication

This manuscript does not contain any individualized data.

### Ethics approval and consent to participate

Not applicable.
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