

Diversity and evolution of telomere and subtelomere DNA sequences in insects

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Abstract

In insects, two types of telomere length maintenance are known: telomerase-dependent, resulting in chromosome ends consisting of short nucleotide repeats (typically TTAGG), and telomerase-independent, resulting in chromosome ends consisting of long nucleotide repeats or transposon-like elements. However, only a few species have been previously studied with regard to their telomere DNA sequences. Here, based on analysis of chromosome-level genome assemblies, I present the data on telomere and subtelomere organization for 180 species from 148 genera, 53 families and 8 orders of insects.

Analysis of these taxa reveals that in fact chromosome ends of most insect species have an intermediate structure and consist of numerous arrays of short telomeric repeats interspersed with telomere-specific non-LTR retrotransposons. An unexpectedly high level of diversity of short telomeric motifs (22 variants ranging in length from 1 to 17 nucleotides) is documented. Telomeres consisting of long repeats (from 173 to 374 bp) are confirmed for flies (the order Diptera) and also found in the orders Odonata and Hymenoptera. The most unusual telomere structure is found in the bee *Lasioglossum lativentre*, in which the chromosomes possess the short telomeric repeat TTAGGTCTGGG at only one end, whereas opposing ends terminate with medium and long repeats.

I conclude that different types of telomere organization and numerous variants of long and short T-containing motifs, including the (T)_n mononucleotide sequence, are compatible with the performance of telomere functions. I argue that both telomerase-dependent and telomerase-independent mechanisms for maintaining telomere length operate simultaneously in many insects. The balance between them and the exchange of sequences between telomeres and subtelomeres are most likely the key factors that determine the structure and evolution of telomeres.

Significance

Multilayer telomeres, resulted from numerous, site-specific insertions of retrotransposons into the region of short telomeric repeats, are not an aberrant type of organization, as previously thought. They are widely distributed among insects and can represent up to 30-40 % of eukaryotic species diversity. Accordingly, the telomere maintenance mechanism based on the joint work of telomerase-dependent and telomerase-independent mechanisms can also be extremely widespread in nature.

Key words: telomere, telomerase, telomeric motif, telomere-specific non-LTR retrotransposons, subtelomere, chromosome-level genome assembly

INTRODUCTION

Telomeres are protein-DNA complexes located at the ends of eukaryotic linear chromosomes (Blackburn, 1991; Osterhage & Friedman, 2009, Nandakumar & Cech, 2013; Shay & Wright, 2019). Telomeres protect the terminal regions of chromosomal DNA from progressive degradation and ensure the integrity of chromosomes (Zakian, 1995; Frydrychová et al., 2021). These essential functions are conserved in eukaryotic organisms and have not changed over hundreds of millions of years of their evolution (Saint-Leandre & Levine, 2020). The persistence of these functions correlates well with the stability in the primary structure of telomeric DNA. In most organisms, as diverse as protozoans, fungi, metazoans, and plants, telomeric DNA consists of the short TTAGGG motif repeated many times (Louis & Vershinin 2005; Červenák et al., 2021). This specific structure of telomere is maintained by telomerase, a telomere-specific reverse transcriptase that elongates the ends of DNA using a small region of its RNA subunit as a template for the telomeric motif (Greider & Blackburn, 1989).

Although telomerase RNA varies dramatically in size, primary sequence and secondary structure among different groups of eukaryotes (Logeswaran et al., 2021), its template part is conservative (Chen et al., 2000; Červenák et al., 2021). Changes in this template are rare in evolution, and as a rule, changed motifs mark higher taxa such as flowering plants (TTTAGGG), nematodes (TTAGGC) and arthropods (TTAGG) (Vítková et al., 2005; Kuznetsova et al., 2020).

However, in some groups of organisms such as alveolate protists (Blackburn & Gall, 1978, Klobutcher et al., 1981; Shirley, 1994; Sohanpal et al., 1995; Liu et al., 1998), ascomycetous yeasts (Červenák et al., 2021) and plant genera *Cestrum* and *Allium* (Peška et al., 2015; Fajkus et al., 2016; Peska & Garcia, 2020), the structure of telomeric motifs changes noticeably faster, and the motifs themselves are more diverse.

Insects are of particular interest for studying the organization and evolution of telomeres, as well as for studying the mechanisms for maintaining telomere length. The telomere motif (TTAGG)_n is considered to be ancestral for the class Insecta and the entire phylum Arthropoda and is preserved in the majority of the studied insect orders and families (Frydrychová et al., 2004; Vítková et al., 2005; Lukhtanov & Kuznetsova, 2010; Kuznetsova et al., 2020). In some beetles (Coleoptera) it is known to be replaced by similar motifs TCAGG and TTAGGG (Prušáková et al., 2021). It can be assumed that the telomere length in these insects is maintained by the standard telomerase-dependent mechanism (Korandová et al., 2014) (see also Mason et al., 2011, 2016). However, for insects of the order Diptera, a completely different, telomerase-independent mechanism has been described, resulting in chromosome ends consisting of long repeats or transposon-like elements (Biessmann & Mason, 1997; Mason et al., 2008). Nematocera species, the so-called lower Diptera, have longer repeated sequences at their chromosome ends that may elongate telomeres by gene conversion (Nielsen & Edstrom, 1993; Rosén & Edström, 2002). *Drosophila* species (higher Diptera, suborder Brachycera) carry terminal retrotransposons that elongate telomeres by transposition specifically to chromosome termini (Biessmann et al., 2000; Casacuberta & Pardue, 2003).

Telomere-specific non-LTR retrotransposons are known to be involved in telomere organization and function in a few other, non-Diptera insects that display the standard TTAGG motif at the ends of chromosomes. In these insects, arrays of the main short motif are interspersed with (or contacted by) retrotransposons (Okazaki et al., 1995; Takahashi et al., 1997; Fujiwara et al., 2005; Garavís et al., 2013). However, the prevalence of this type of telomere organization and its possible role in maintaining telomere length are poorly understood (but see Garavís et al., 2013).

Another problem in the study of insect telomeres is the so-called “loss of the ancestral TTAGG motif”, which has been repeatedly reported for members of several insect orders (Frydrychova & Marec, 2002; Mason et al., 2016; Kuznetsova et al., 2020; Prušáková et al., 2021), for example, for several families of Hemiptera (Frydrychová et al., 2004; Golub et al., 2015, 2017, 2018), Hymenoptera (Menezes et al., 2017; Gokhman & Kuznetsova, 2018) and Odonata (Kuznetsova et al., 2021). Reports of the loss of the TTAGG motif are based on numerous studies in which TTAGG (or similar motifs) were searched using fluorescent in situ hybridization (FISH) and/or southern blot hybridization techniques. Both techniques can detect a specific DNA sequence, but unfortunately are not available for detecting DNA fragment with unknown DNA sequence (Grozeva et al., 2011). Therefore, the true reasons for this “loss of the ancestral TTAGG motif” (transition to another short motif or a complete change in the structure of telomeres, for example, replacement by long repeats) has remained unknown.

Here, based on analysis of chromosome-level genome assemblies, I present the data on telomere and subtelomere organization for 180 species from 148 genera, 53 families and 8 orders of insects.

This approach, based on the analysis of nucleotide sequences of whole chromosomes, solves the problem of the “lost TTAGG motif”, reveals the true diversity of telomeric and subtelomeric organization, and outlines approaches to solving the question of how insect telomeres evolved.

MATERIAL AND METHODS

Chromosome-level genome assemblies generated by the Darwin Tree of Life Project (<https://www.darwintreeoflife.org/>) (The Darwin Tree of Life Project Consortium, 2022) and freely available upon deposition in the European Nucleotide Archive (ENA) (<https://www.darwintreeoflife.org/wp-content/uploads/2020/03/DToL-Open-Data-Release-Policy-1.pdf>) were used for analysis of telomere and subtelomere sequences (Table S1). Published and freely available chromosome-level genome assemblies of *Formica selysi* (Brelsford et al., 2020), *Solenopsis invicta* (Yan et al., 2020), *Nasonia vitripennis* (Benetta et al., 2020) and *Apis mellifera* (Wallberg et al., 2019) were also used.

The chromosome-scale scaffolds confirmed by the Hi-C data were analyzed using the graphical sequence panel (format “Graphics”) implemented in GenBank. Alternatively, in some cases, the chromosome-scale scaffolds were downloaded and analyzed in BioEdit (Hall, 1999). The terminal regions of chromosomes were first visually inspected. Then tandem repeats were searched using Tandem Repeats Finder (Benson, 1999). Finally, the position analysis and the comparison of putative telomeric and subtelomeric repeats were performed using the “Search” option in the BioEdit program and the “Search” and “BLAST” options implemented in GenBank.

In almost all species in which a distinct short telomere motif has been found in the majority of chromosomes, in some chromosomes the telomere motif was absent on one or even both chromosome ends. This situation has been previously described for the honey bee (Wallberg et al., 2019). As for the honey bee, I interpret these chromosome-scale scaffolds as truncated ones, but not as lacking arrays of telomere motifs. A telomere was considered present if its length exceeded 400 bp, the estimated minimal functional length required for t-loop formation (Watson et al., 2021).

The search of the telomere-specific non-LTR retrotransposons was conducted manually (Goubert et al., 2022) in 500000 bp terminal fragments of the 5' ("left") and 3' ("right") chromosome ends, by using the contact of the polyA (polyT) tract with short telomeric repeats (TTAGG or others) as a specific marker (Okazaki et al., 1995; Takahashi et al., 1997).

In accordance with common practice, hereinafter in the text (except Figure 2) I list telomeric and subtelomeric sequences as if they were found at the 3' chromosome end ("right" telomere), even if in Genbank they are given for the 5' chromosome end ("left" telomere) and have, respectively, a reverse complement orientation and spelling.

RESULTS AND DISCUSSION

Repeats forming the extreme ends of chromosomes (telomeric motifs)

In the studied species, 22 variants of the main short telomeric motifs were found at the very ends of chromosomes: 1bp repeat (T), 5 bp repeats (TTAGG, TCAGG, TTGGG), 6bp repeat (TTAGGG, TTCCTC, GGGTCT), 8 bp repeat (TTATTGGG), 9 bp repeat (TGGATGGGA), 10 bp repeats (TTAAAACGCC, TTGAAACGCC, TTAAGCGCC, TTAAGGCGTT, TTAGGGTGGT, TTAGTTTGGG), 11 bp repeats (TTAGGTCTGGG, TTAGGTTGGGG, TTAGGTTCCGGG, TTGGGTCTGGG, TTGCGTCTGGG, TTGCGTCAGGG) and 17 bp repeat AAAAAATTCTTTGATGC (Table S1). Of these motifs, three were previously known for insects (TTAGG, TCAGG, TTAGGG) (Frydrychová et al., 2004; Kuznetsova et al., 2020; Prušáková et al. 2021), and 19 are described here for the first time.

It is not surprising that the 5 bp motif TTAGG, "standard" for insects (Frydrychová et al., 2004; Kuznetsova et al., 2020) was found at the ends of chromosomes in most of the studied species (Table S1). Confirming already known data (Frydrychová et al., 2004; Mandrioli et al., 2012; Vershinina et al., 2015), this motif was found to be invariable within the orders Lepidoptera and Trichoptera (butterflies and moths, caddisflies). It was also found in the only studied species of the order Plecoptera, in most beetles, in one of the basal lineages of the order Hymenoptera (Tenthredinidae: *Tenthredo notha*) and in several representatives of the clade Aculeata (Hymenoptera) (Fig. 1). In beetles (the order Coleoptera), the modified 5 bp motif TCAGG was found in two species (Melyridae: *Malachius bipustulatus*; Pyrochroidae: *Pyrochroa serraticornis*), and the modified 5 bp motif TTGGG was found in the species *Apoderus coryli* (Atelabidae). The motif TTGGG was also found in the white-legged damselfly *Platycnemis pennipes* (Odonata, suborder Zygoptera: Platycnemididae).

Strongly modified 10 bp motifs were found in pentatomomorphic bugs *Aelia acuminata* (Hemiptera, Pentatomidae) (TTAGGGATGG) and *Acanthosoma haemorrhoidale* (Hemiptera,

Acanthosomatidae) (TTAGGGTGGT). (The loss of the TTAGG motif for the infraorder Pentatomomorpha was previously reported (Grozeva et al., 2011)).

The most intriguing results were obtained in the study of the order Hymenoptera (Fig. 1, Table S1). In the “primitive” family Tenthredinidae, which represents one of the first branches within the Hymenoptera, a common sawfly *Tenthredo notha* had the TTAGG motif, ancestral to all arthropods. This result is consistent with previously obtained, FISH-based data on two other species of this family (Gokhman & Kuznetsova, 2018).

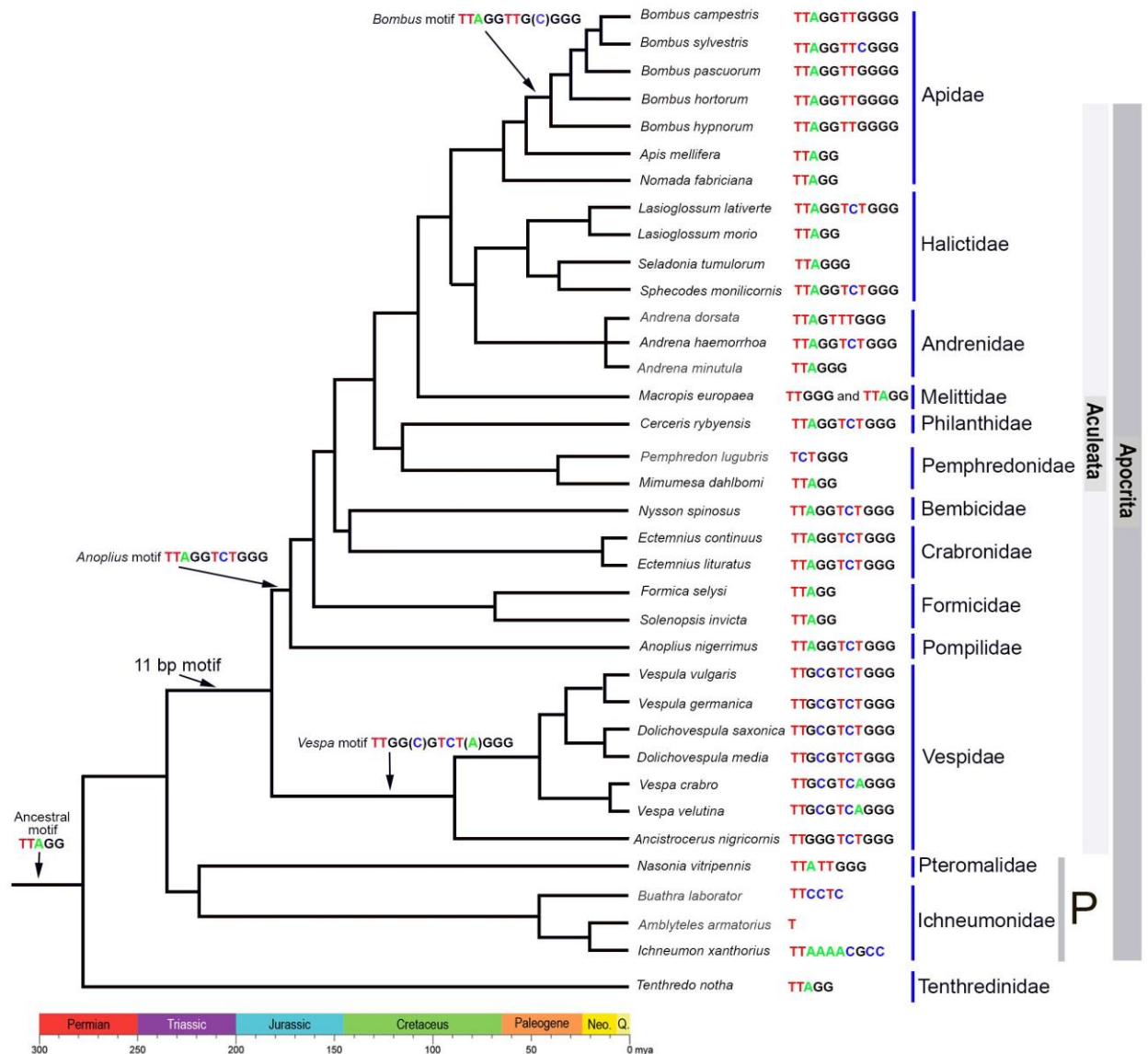


Figure 1. Diversity and evolution of short telomeric motifs in the order Hymenoptera. The phylogeny and dating are based on the published works (Cameron et al., 2007; Peters et al., 2017; Sann et al., 2018, 2021). P is Parasitica (Here: Pteromalidae + Ichneumonidae).

The clade Apocrita showed a clear trend to a high diversity of short telomeric motifs, with the prevalence of multi-nucleotide motifs (8-11 bp). In these multi-nucleotide motifs, the first five letters were often represented by the classic TTAGG (or slightly modified variants), while the G and T nucleotides predominated in the second part of the motifs.

In four studied species of parasitic wasps (Parasitica, a group of the infraorder Apocrita), the 5 bp TTAGG motif was not found (Table S1, Fig. 1), confirming the previous results of Gokhman et al. (2014). Instead, 6 bp motif TTCCTC was found in *Buathra laborator* (Ichneumonidae) and various motifs, with predominance of 10 bp motif TTAACGCGCC and the 8 bp motif TTATTGGG were found in *Ichneumon xanthorius* (Ichneumonidae) and *Nasonia vitripennis* (Pteromalidae) respectively.

The most unusual telomere structure was found in the parasitic wasp *Amblyteles armatorius* (Ichneumonidae). In this species a mononucleotide telomeric motif T was found. It consisted of the (T)_n sequence at the 3' chromosome end and (A)_n sequence at the 5' chromosome. The length of this mononucleotide telomere ranged from 306 to 629 bp, being comparable with the estimated minimal functional length required for t-loop formation (Watson et al., 2021).

The clade Aculeata, the group containing stinging wasps, ants and bees, was characterized by the presence of 11 nucleotide motifs. The TTAGGTCTGGG motif predominates in different families and is probably ancestral for the Aculeata.

Groups of specific motifs, that are derived from TTAGGTCTGGG, were found in the family Vespidae (TTGGGTCTGGG, TTGCGTCAGGG, TTGCGTCTGGG) and in the bumble bees (genus *Bombus*) (TTAGGTTGGGG, TTAGGTTGGG).

The 6 bp motifs were found in the bee *Seladonia tumulorum* (Halictidae) (TTAGGG) and the wasp *Pemphredon lugubris* (Pemphredonidae) (TCTGGG).

The 10 bp motif TTAGTTTGGG was found in the bee *Andrena dorsata* (Andrenidae).

In addition, six cases of possible reversion to the ancestral TTAGG motif were identified in Aculeata: (1) in the family Formicidae, (2) in *Mimumesa dahlbomi* (Pemphredonidae), (3) in *Macropis europea* (Melittidae), (4) in *Lasioglossum morio* (Halictidae), (5) in *Nomada fabriciana* and (6) *Apis mellifera* (Apidae). Alternatively, we can think that the TTAGG motif was ancestral for Aculeata; then we must assume the multiple cases of independent origin of the TTAGGTCTGGG motif in different branches of Aculeata.

Short telomeric repeats were not found in flies (Diptera), which is also in line with expectations (Rosén & Edström, 2002; Mason et al., 2008; Kuznetsova et al., 2020). Instead, in five species of the suborder Brachycera, long repeats were found at the ends of chromosomes. The length of these repeats varied from 173 to 374 bp. Alignment of these long telomeric repeats from different ends of the same chromosome, from different chromosomes of the same species, and from different species showed: (1) species specificity and small variations (95-98% similarity) in length and nucleotide composition within each species, and (2) sharp interspecific differences, while maintaining some similarity between the species, which most likely indicates the origin of these repeats from a common ancestor.

Short telomeric repeats were not found in the blue-tailed damselfly *Ischnura elegans* (Odonata, Zygoptera: Coenagrionidae). In this species, various sequences, including low-copy repeats of various length, were found at the very ends of autosomes, while 175 bp long repeats formed the very terminal regions of the X chromosome.

Intraspecific stability and variation of short telomere motifs

The length of the most terminal telomeric sequences in the studied species varied from 365 to 15724 bp and included from 33 to 1989 short repeats. Within this length, the structure of the main motifs was stable in most cases, sometimes with rare single altered variants, which can be considered as single substitutions/indels or sequencing errors.

However, in the order Hymenoptera, several species were found in which the main motifs were regularly interspersed with variant repeats.

In the bumble bee *Bombus campestris*, the main repeat 11 bp motif TTAGGTTGGGG was found to be interspersed with the sequences TTAGGTGGGGG, TTAGGTTGGTGGGG, TTAGGTTGGTTAGTGGGG, TTAGGTTGGGTAGGTTTGG, TTAGGTTGGGT and TTTGGGTTGGGG. In the bumble bee *B. sylvestris*, the main 11 bp repeat TTAGGTTTCGGG was found to be regularly interspersed with 16 bp motif TTAGGTTAGGTTTCGGG.

In the pteromalid wasp *Nasonia vitripennis*, the main motif TTATTGGG obviously prevailed, but the modified sequences TTATTGGGGG, TTTTATTGGG, TTTTATTGGG, TTATTGG, TTATTGGAGGG, TTATTGGGG, TTATTATTGGG and TTTTGTTATTGGG were regularly found.

Only partially confirming previous analyses (Garavís et al., 2013; Wallberg et al., 2019), in the honey bee *Apis mellifera*, the TTAGG telomeres were found to be interspersed with variant repeats TTAGGGTT and TTATAGG.

In the bee *Macropis europaea*, the two main motives TTGGG and TTAGG were found.

In the beetle *Philonthus cognatus* the main motif TTAGG interspersed with TTTAGG.

In the parasitic wasp *Ichneumon xanthorius* the motif TTAAAACGCC was found to be fixed in chromosome 11 and to prevail in chromosomes 1 and 8 and in the left telomere of chromosome 2, in which it was interspersed with variant motives TTAAAACGCCT, TTAAACGCCT, TTACAACGCC, TTAGAACGCC, TTAAACCGCC, TTACAACGCC, TTACGACGCC and TTAAATCGCC. In chromosome 6, the motif TTAAAACGCC and the altered motif TTGAAACGCC occurred with approximately equal frequency. The motif TTAAAGCGCC was fixed in chromosome 9. In chromosome 7, the strongly altered motif AAAAAATTCTTTGATGC occurred. In chromosome 5, the strongly altered main motif TGGATGGGA was found to be interspersed with variant sequences CGGGA, TGGGA and TGGGA. Finally, in the right telomere of chromosome 2, the main motif TTAAGGCGTT (which is actually a palindrome of the main motif in the left telomere!) was interspersed with variant repeats TTAAAGCCGTT, TTGTAAGGCG, TTCTAAAGGCG and TTAAAGGCGTT.

The most unusual telomere structure and intraspecific variations were found in chromosomes 1-13 of the bee *Lasioglossum lativentre*. In these chromosomes, one end is formed by arrays of a short motif TTAGGTCTGGG associated with subtelomeric sequence (TTAGGTCTGCATCGCGG)_n or *SART* elements, and the other end is formed by medium and long repeats (similar to Diptera) (Table 1). The structure of the first ends (without the TTAGGTCTGGG motif) is difficult to explain by truncation, since they do not contain not only the telomeric TTAGGTCTGGG motifs, but also the specific subtelomeric sequences of the second ends (although independent confirmation of telomere structure in this unusual case would be desirable, for example, using the FISH method to reveal the structure of telomeres on microscopic preparations). Interestingly, a similar structure was recently discovered in the tapeworm

Hymenolepis microstoma (Platyhelminthes), in which the chromosomes possess telomeric repeats GGGATT at only one end, whereas opposing ends terminate with a novel repeat array with a median unit length of 179bp (Olson et al., 2020).

Table 1. Telomere and subtelomere sequences in chromosomes of the bee *Lasioglossum lativentre*.

Chromosome #	First chromosome end: telomere	First end: subtelomere	Second chromosome end: telomere
1	TTAGGTCTGGG	<i>SART</i> -(TTAGGTCTGGG) _n	41 bp repeats
2	TTAGGTCTGGG	<i>SART</i> -(TTAGGTCTGGG) _n	304 bp repeats
3	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	150 bp repeats
4	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	118 bp repeats
5	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	304 bp repeats
6	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	Various short, medium and long repeats
7	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	Various short, medium and long repeats
8	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	Various short, medium and long repeats
9	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	40-41 bp repeats
10	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	41 bp repeats
11	TTAGGTCTGGG	<i>SART</i> -(TTAGGTCTGGG) _n	211 bp repeats
12	TTAGGTCTGGG	(TTAGGTCTGCATCGCGG) _n	Various short, medium and long repeats
13	TTAGGTCTGGG	<i>SART</i> -(TTAGGTCTGGG) _n	148 bp repeats
14	41 bp repeats		41 bp repeats

Sequences adjacent to the most terminal telomeric repeats (Telomere-Subtelomere structure)

Subtelomeric DNA consists of specific repetitive sequences that are in direct contact with telomeres and, accordingly, have a subterminal position in chromosomes (Louis & Becker, 2014). As a rule, subtelomeres are more diverse in structure of their DNA sequences, which is correlated with the diversity of subtelomeric proteins (Louis & Vershinin, 2005; Saint-Leandre & Levine, 2020).

In order to study the diversity of the subtelomeric region, in the species under study I inspected the sequences that were adjacent to the most terminal chromosomal array. This search revealed three types of the subterminal organization. The sequences of the first two types had

features typical of subtelomeres (Louis & Becker, 2014), while the third type is represented by multilayer telomeres that share some of the properties of subtelomeres (see below).

(Type 1) The subtelomeric region of the first type was found to include numerous tandem repeats of various lengths. This type was found in the caddisfly *Limnephilus lunatus* (Trichoptera), the bug *Aelia acuminata* (Hemiptera), in representatives of the orders Hymenoptera (*Anoplius nigerrimus*, *Bombus sylvestris*, *Ectemnius continuus*, *Nasonia vitripennis*, *Nomada fabriciana*, *Nysson spinosus*, *Seladonia tumulorum*, *Tenthredo notha*; part of chromosomes of *Andrena haemorrhoea*, *Lasioglossum morio* and *Mimumesa dahlbomi*) and Coleoptera (part of chromosomes of *Apoderus coryli*) (Table S1).

A variant of the first type were subtelomeres, which consisted only of long repeats. This variant was found in two species: in the right telomere of chromosome 6 in the beetle *Apoderus coryli* (100 repeats of the 171 bp motif) and in the bee *Macropis europaea*, in which the subtelomere consisted predominantly of 121 bp repeats (Table S1). This variant was previously reported for the aphid *Myzus persicae* (Hemiptera) (Spence et al., 1997). Such subtelomeres, consisting of long repeats, are structurally very similar to the telomeres found by me in five species of flies (Diptera: suborder Brachycera) (Table S1) and previously in *Chironomus* (Diptera: suborder Nematocera) (Rosén & Edström, 2002).

(Type 2) The second type, which is widespread in the order Hymenoptera (the families Apidae, Crabronidae, Formicidae, Halictidae, Philanthidae, Vespidae), was represented by subtelomeres consisting of modified telomeric sequences (Table S1). These modified sequences were found to include: (a) altered versions of the main motif; (b) elongated and varying in structure variants that retained similarities with the main motif, and (c) longer repeats (20-41 bp) that had a remote resemblance to the main motif. Interestingly, these longer repeats may have high specificity. For example, the 41 bp subtelomeric repeat AAAATACCTTATAATATAACGTAATATGCAGTGTATCATTG was detected in all 14 chromosomes of the bee *Lasioglossum lativentre*, but the BLAST function throughout the Genebank database did not reveal it in any other organisms. Кроме того, в субтеломерном региона могут встречаться отдельные копии неизмененного основного теломерного мотива. In addition, copies of the unaltered main telomeric motif may sporadically occur in the subtelomeric region, for example, in the wasp *Dolichovespula sylvestris*.

(Type 3) The most widespread among the studied species and the most intriguing was the third type of subterminal organization, which was found to be characterized by numerous arrays of short repeats (for example, TTAGG in Lepidoptera, TTAGGTTGGGG in bumble bees) interspersed with telomeric repeat-specific non-LTR retrotransposons of the *SART* and *TRAS* families.

TRAS is a family of retrotransposable, telomeric repeat-associated elements with a polyA (polyT) tail at the end (Okazaki et al., 1995). It is integrated in a highly specific manner into the short telomeric repeats at the same target sequence. In right telomere (3' chromosome end) it can be easily recognized by its typical long (14-72 (T)_n) fragment, immediately starting after the GG dinucleotide of the short repeat (Fig. 2).

SART is a family of retrotransposable, telomeric repeat-associated elements with a polyA (polyT) tail at the end, in a reverse orientation from that of *TRAS* (Takahashi et al., 1997). It is inserted into the telomeric repeats in a highly specific manner. Therefore, in right telomere (3' chromosome end) it can be easily recognized by its typical long (14-72 (A)_n) fragment contacting with the nucleotides GG(GGG) and then followed by standard short motifs (Figs 2 and 3).

For the silkworm *Bombyx mori*, the retrotransposons *TRAS* and *SART* are sometimes considered as subtelomeric elements because they do not occupy the most terminal position, being proximal to the most distal array of canonical short repeats (Kubo et al., 2001; Fujiwara, 2014; Louis & Becker, 2014; Mason et al., 2016). In fact, they are an integral part of the telomere, since they are inserted specifically and exclusively into telomeric repeats and cannot exist outside of short telomeric motifs (Osanai et al., 2006).

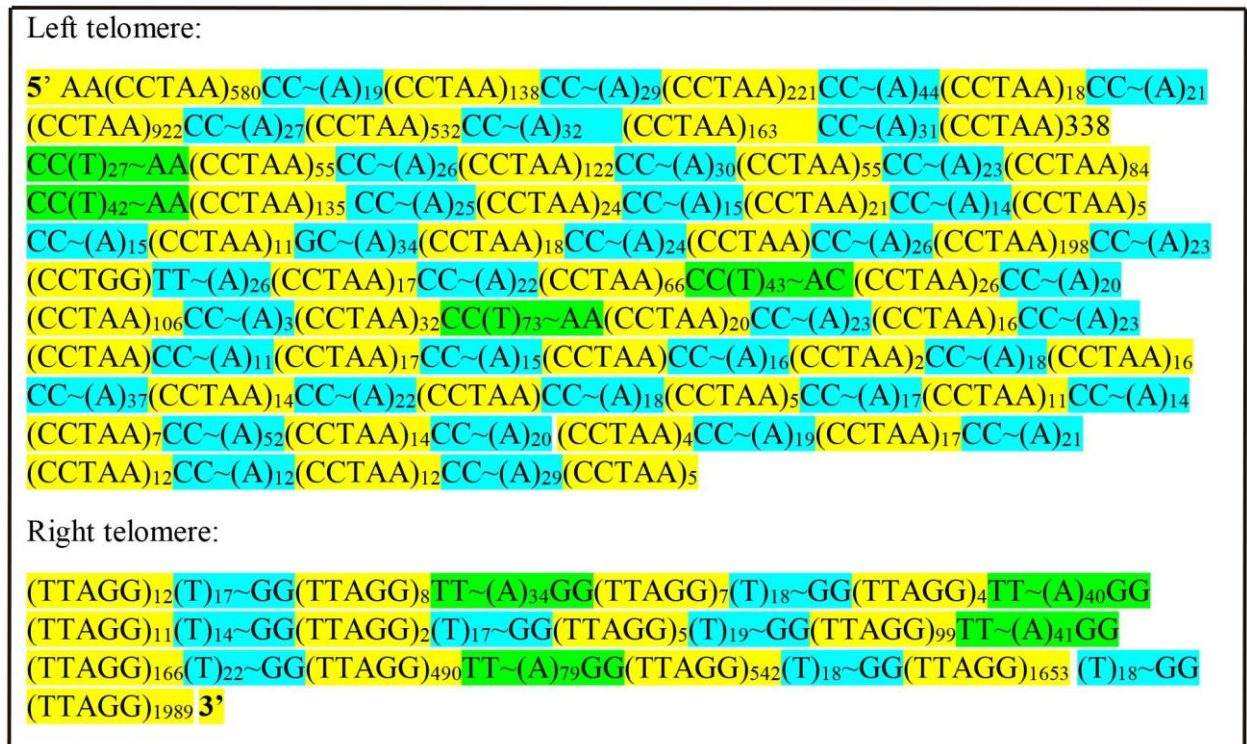


Figure 2. Telomere structure in chromosome 1 of *Orgyia antiqua* (Lepidoptera).

Short telomeric repeats (CCTAA in left (5') and TTAGG in right (3') chromosome ends) (yellow) are interspersed with retrotransposable elements *TRAS* (blue) and *SART* (green). Left telomere (positions from 1 to 266133) includes 44 arrays of TTAGG interspersed with 39 *TRAS* and 4 *SART* retrotransposons. Right telomere (positions from 44592109 to 44675926, 83817 bp) includes 13 arrays of TTAGG interspersed with 8 *TRAS* and 4 *SART* retrotransposons. The tilde symbol (~) denotes the main body of *TRAS* and *SART* (without polyA/polyT tail); its length varies from 614 to 15369 bp, with a clear predominance of sequences from 6000 to 7000 bp.

To the best of my knowledge, this type of organization has previously been reported for only six species of insects: three species of moths (Kubo et al., 2001); flour beetle *Tribolium castaneum* (Osanai et al., 2006), pea aphid *Acyrtosiphon pisum* (Hemiptera) (International Aphid Genomics Consortium, 2010), human body louse *Pediculus humanus humanus* (Phthiraptera) (Kirkness et al., 2010). It was also known for the mite *Tetranychus urticae* (Grbić et al., 2011).

Here it was found in 113 species of Lepidoptera, 10 species of Coleoptera, 10 species of Hymenoptera, 3 species of Trichoptera and 1 species of Hemiptera (Table S1). The number of retrotransposons inserted in telomeric regions was found to vary from single ones found in some (not all) chromosomes (for example, in *Mimusesa dahlbomi* only in chromosome 9, in *Coccinella septempunctata* only in chromosome 8) to several tens at each chromosome end. In the latter case,

the massive integration of these elements into the proximal regions of the TTAGG, TTGGG, TCAGG, TTAGGTTGGGG, TTAGGTCTGGG, and TTGGGTCTGGG repeat arrays resulted in huge telomeric massive up to 260 kb in size (Fig. 2).

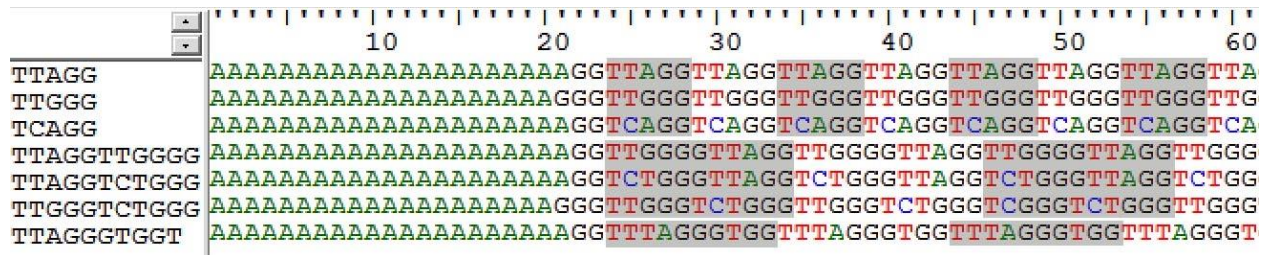


Figure 3. Contact area between the (A)_n-tail of the *SART* retrotransposon and short telomeric repeats TTAGG, TTGGG, TCAGG, TTAGGTTGGGG, TTAGGTCTGGG, TTGGGTCTGGG, and TTAGGGTGGT.

The telomeric repeat-specific non-LTR retrotransposons seems to be highly species-specific. For example, in *Bombus hortorum*, one of the retrotransposons of the *SART* family has a length of 5951 bp. It was found in all 18 chromosomes of this species, and the similarity between chromosomes was 99-100%. The BLAST function revealed this (or similar) element in *Bombus vancouverensis*, *B. bifarius*, *B. vosnesenskii*, *B. sylvestris*, *B. campestris*, *B. terrestris*, *B. pasuorum*, as well as in all 12 chromosomes of *B. hypnorum*, but the similarity between different species was in the range of 69-80%.

Interestingly, the three types of subterminal organization of chromosomes found are not necessarily fixed within a species or even individual chromosomes (Table S1); for example, the beetle *Apoderus coryli* has telomeres of all three of these types. In general, subtelomeric sequences are much more variable than telomeric sequences.

CONCLUSION

This study solves the problem of the lost of the TTAGG telomeric motif in insects (Frydrychova & Marec, 2002; Mason et al., 2016; Kuznetsova et al., 2020, 2021; Prušáková et al., 2021) and shows that the absence of this motif in representatives of the orders Hemiptera and Hymenoptera is explained by evolutionary transitions to other, previously unknown short telomeric repeats. In the order Odonata (suborder Zygoptera), the absence of the TTAGG motif (Kuznetsova et al., 2021) can be explained by both nucleotide substitutions (as it is observed in *Platycnemis pennipes*, which have TTGGG instead of the ancestral TTGGG motif) and by a complete change in the telomere structure due to loss of short telomeric repeats (as seen in *Ischnura elegans*).

The study revealed an extremely high diversity in the structure of telomeric nucleotide sequences in insects. This diversity is manifested in:

- (1) numerous (22) variants of short telomeric motifs found in different orders, especially in Hymenoptera (Table S1),
- (2) in presence of telomeres consisting of long repeats, especially in Diptera, but also in Odonata and Hymenoptera (Table S1), and
- (3) different types of intra-telomeric organization (Fig. 4).

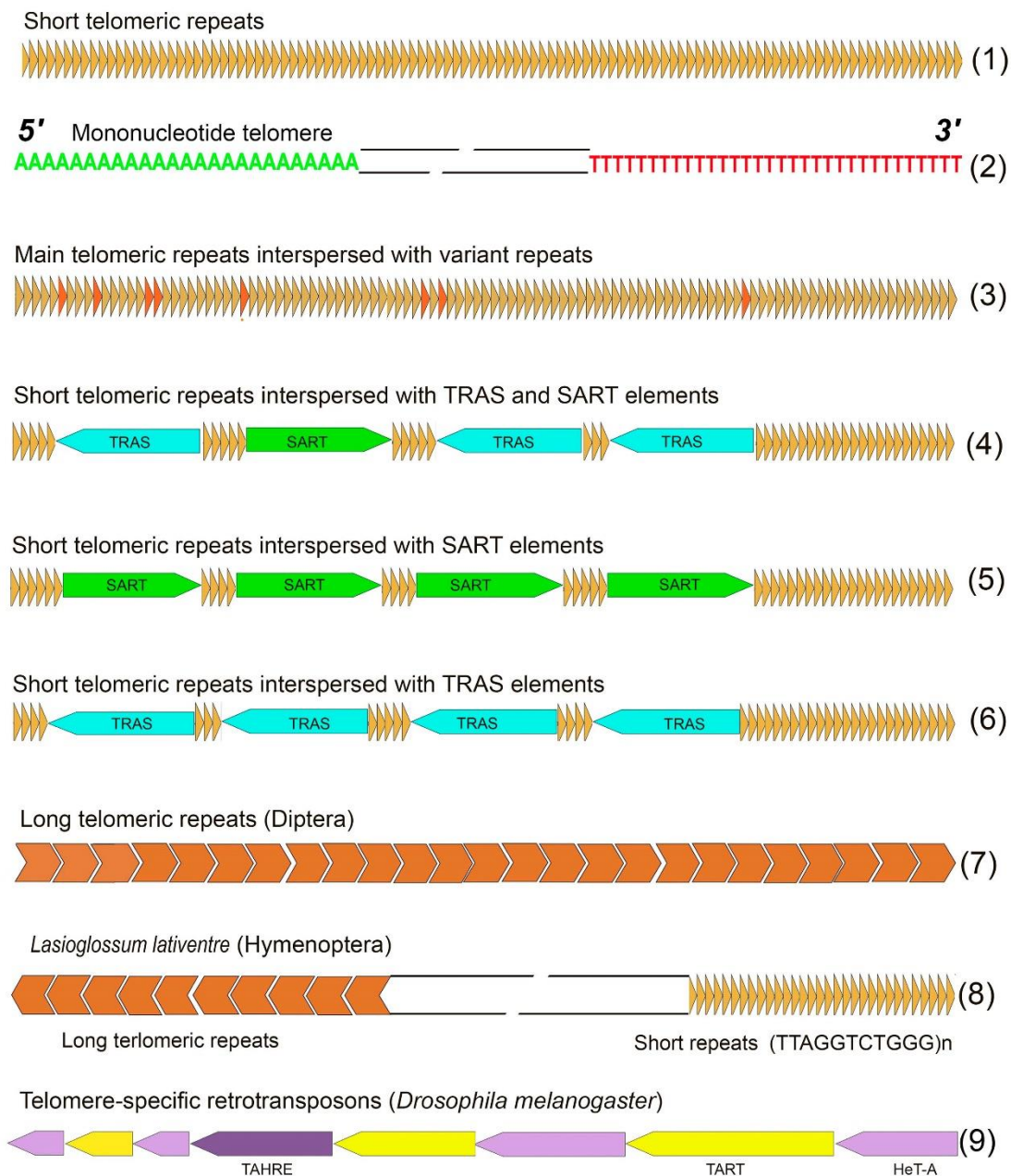


Figure 4. Types of telomere structures and telomere sequences in insects.

(1) short (5-17 bp) repeats (canonical telomere); (2) mononucleotide telomere, consisting of long (306-629 bp) (A)_n tail at the 5' chromosome end and (T)_n tail at the 3' chromosome end (*Amblyteles armatorius*, Ichneumonidae); (3) main short repeats interspersed with variant short repeats; (4) main short repeats interspersed with telomere-specific non-LTR retrotransposons of the *TRAS* and *SART* families; (5) main short repeats interspersed with telomere-specific non-LTR retrotransposons of the *SART* family; (6) main short repeats interspersed with telomere-specific non-LTR retrotransposons of the *TRAS* family; (7) long (173-374 bp) repeats, (8) one telomere consists of long repeats (similar to Diptera) and the other telomere if formed by arrays of short repeats TTAGGTCTGGG; (9) telomere-specific non-LTR retrotransposons of the *HeT-A*, *TAHRE* and *TART* families.

The study confirms that the pentanucleotide TTAGG motif is widespread among insects. However, telomeres consisting solely of the TTAGG repeats (type 1 in Fig. 4) turned out to be not as common (7 out of 180 species studied) as it is previously believed (Frydrychová et al., 2004; Vítková et al., 2005; Lukhtanov & Kuznetsova, 2010; Kuznetsova et al., 2020). Other types dominate or are found in six insect orders (Fig. 4): type 6 in Trichoptera, type 4 in Lepidoptera, type 5 in Coleoptera and Hemiptera, types 3 and 5 in Hymenoptera, and types 7 and 9 in Diptera. Interestingly, the last five orders are the richest in species number. Together they comprise over 800,000 described species, i.e. about 80% of the known insect species richness (Stork, 2018). The “non-canonical” telomeric organizations actually characterize a significant part (~40%) of all *ca*1900,000 described eukaryotic species (Chapman, 2009).

The most unexpected telomere sequence, consisting exclusively of the nucleotide T at the 3' chromosome end, was found in the parasitic wasp *Amblyteles armatorius* (type 2 in Fig. 4). A common feature of all identified telomeric motifs (Table S1, Fig. 1) is the predominance of the nucleotide T. Telomeric motifs are generally thought to be saturated with G nucleotide (Osterhage & Friedman, 2009; Garavís et al., 2013); however, telomeric motifs without guanine (TTCCTC and T) were found in two parasitic wasp species (Fig. 1). Thus, different types of telomere organization and different variants of T-containing short motifs, including mononucleotide sequences, are compatible with the performance of telomere functions.

Therefore, the length and nucleotide composition of short telomeric motifs in themselves do not appear to be significant constraints in the evolution of telomeres. In this situation, when there is no mandatory selection for only one type of motif, and if telomerase is lost, as suggested by some theories (Mason et al., 2016), one would expect rapid evolution of telomere nucleotide sequences as a result of genetic drift/selection. However, this expectation is generally not met. In fact, the ancestral motif TTAGG is preserved in most arthropods over hundreds of millions of years of their evolution (Vítková et al., 2005; Kuznetsova et al., 2020).

Altered variants of telomeric motifs are also phylogenetically stable, although to a lesser degree. For example, in the order Hymenoptera, the modified 11 bp motifs characterize high-level clades, families, genera, and groups of species (Fig. 1). Such evolutionary stability of the motifs is easily explained by their maintenance through the telomerase-dependent mechanism: the telomeric template region of the telomerase RNA subunit is conservative (Chen et al., 2000; Červenák et al., 2021); therefore, its changes are extremely rare, and if they occur, they mark higher taxa.

On the other hand, in some Hymenoptera species, the detected high intra-telomeric and, even more so, inter-telomeric and inter-chromosomal variability, is hardly compatible with the telomerase-dependent mechanism of telomere length maintenance. It is hard to imagine that within the same organism there are many differentiated RNA templates that support the polymorphism of telomeric motifs. Such intraspecific variations seem to be more compatible with traditional Darwinian evolution, in which new motifs arise from nucleotide substitutions and then propagate and become fixed in individual chromosomes and genomes through genetic drift and/or selection.

Site-specific insertions of the *SART* and *TRAS*-retrotransposons into the telomeric region is found in most of the studied species, leading to the formation of multilayer telomeres (types 4-6 in Fig. 4) in five of eight studied orders. This variant of the “non-canonical” telomeric organization is very common in the order Hymenoptera, prevails in the order Coleoptera and is the only type in the order Lepidoptera. The prevalence of this “non-canonical” telomeric organization

in other insect orders remains to be studied, but it can be assumed to be common, given that it is already found in the orders (Hemiptera) (International Aphid Genomics Consortium, 2010) and (Phthiraptera) (Kirkness et al., 2010) and also outside the class Insecta in the mite *Tetranychus urticae* (Grbić et al., 2011). Interestingly, the multilayer telomeric organization correlates with low telomerase activity and loss of introns and GQ domains from TERT genes (Fujiwara, 2014). Most likely, the telomere length in these insects is partially maintained through transpositions and terminal recombination (Fujiwara et al., 2005; Osanai et al., 2006).

However, a pure *SART/TRAS*-based telomere maintenance mechanism seems impossible without contribution of the telomerase-based processes. The reason is that retrotransposons can integrate into telomeric regions in a highly specific manner. Therefore, if telomere motifs disappear (which is inevitable if telomerase activity is absent, due to shortening of telomeres in a series of cell divisions), this will also lead to the disappearance of the telomere-specific retrotransposons *SART* and *TRAS* (Fujiwara et al., 2005).

Single copies of unaltered or modified telomeric motifs are often found in subtelomeres. I hypothesize that the telomeric repeats can enter the subtelomeric region due to recombination and be modified there due to nucleotide substitutions. Through recombination, these modified repeats can re-enter telomeric regions, resulting in the structure, in which main short repeats interspersed with variant short repeats (type 3 in Fig. 4), as seen in some Hymenoptera.

With a complete loss of telomerase activity, there should be a loss of short telomeric motifs, and then, inevitably, a loss of the *SART* and *TRAS* elements. The intermediate stage of this process is observed in the bee *Lasioglossum lativentre* (loss of part of the telomeres and their replacement by subtelomeres, type 8 in Fig. 4), and the final stage is observed in Diptera (type 7 in Fig. 4).

Thus, the balance between telomerase-dependent and telomerase independent mechanisms and the exchange of sequences between telomeres and subtelomeres are most likely the key factors that determine the structure and evolution of telomeres in insects.

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DATA AVAILABILITY

All the analyzed chromosome-level assemblies are available via the GenBank links provided (Table S1).

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Appendix

Table S1. Telomeric motifs and other sequences found in telomeric and subtelomeric regions in chromosomes of the studied insect species.

Order, Family	Species	Telomere motif (repeats found at the most terminal positions of chromosome ends)	Sequences adjacent to the most terminal chromosomal array	GenBank Accession #
ODONATA				
Zygotera: Platycnemididae	<i>Platycnemis pennipes</i>	TTGGG	Combination of various repeats with prevalence of 129, 286-289 and 434 bp long repeats	OW121847.1- OW121846.1
Zygotera: Coenagrionidae	<i>Ischnura elegans</i>	There is no short telomeric motif. Various sequences, including low-copy repeats of various length in autosomes. 175 bp repeats in X chromosome.s		OV121100.1-OV121106.1
PLECOPTERA				
Nemouridae	<i>Nemoura dubitans</i>	TTAGG	Various sequences including short repeat AACGG	OV121074.1- OV121073.1
HEMIPTERA				
Pentatomidae	<i>Aelia acuminata</i>	TTAGGGATGG	Combination of various short, medium and long repeats	OU426978.1-OU426985.1
Acanthosomatidae	<i>Acanthosoma haemorrhoidale</i>	TTAGGGTGGT	Combination of various short, medium and long repeats or SART - (TTAGGGTGGT) _n	OV884011.1- OV884010.1
LEPIDOPTERA				
Blastobasidae	<i>Blastobasis adustella</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU026114.1-OU026144.1
Blastobasidae	<i>Blastobasis lacticolella</i>	TTAGG	SART - (TTAGG) _n - TRAS	LR990039.1-LR990069.1
Cossidae	<i>Zeuzera pyrina</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU015617.1-OU015648.1
Crambidae	<i>Chrysoteuchia culmella</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342641.1-OU342672.1
Crambidae	<i>Paraponyx stratiotata</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342457.1-OU342487.1
Drepanidae	<i>Habrosyne pyritoides</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU015585.1-OU015616.1
Drepanidae	<i>Thyatira batis</i>	TTAGG	SART - (TTAGG) _n - TRAS	LR990485.1-LR990516.1
Erebidae	<i>Eilema depressum</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU612012.1-OU612042.1

Erebidae	<i>Eilema sororculum</i>	TTAGG	SART - (TTAGG)n - TRAS	OU618532.1-OU618562.1
Erebidae	<i>Euproctis similis</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990103.1-LR990125.1
Erebidae	<i>Hypena proboscidalis</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990126.1-LR990157.1
Erebidae	<i>Laspeyria flexula</i>	TTAGG	SART - (TTAGG)n - TRAS	LR989949.1-LR989980.1
Erebidae	<i>Lymantria monacha</i>	TTAGG	SART - (TTAGG)n - TRAS	LR991081.1-LR991109.1
Erebidae	<i>Orgyia antiqua</i>	TTAGG	SART - (TTAGG)n - TRAS	OU779861.1-OU779875.1
Erebidae	<i>Schrankia costaestrigalis</i>	TTAGG	SART - (TTAGG)n - TRAS	FR997825.1-FR997856.1
Erebidae	<i>Spilarctia lutea</i>	TTAGG	SART - (TTAGG)n - TRAS	OU696470.1-OU696502.1
Erebidae	<i>Spilosoma lubricipeda</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992274.1-HG992304.1
Geometridae	<i>Agriopsis aurantiaria</i>	TTAGG	SART - (TTAGG)n - TRAS	OU611981.1-OU612011.1
Geometridae	<i>Biston betularia</i>	TTAGG	SART - (TTAGG)n - TRAS	FR989862.1-FR989893.1
Geometridae	<i>Campaea margaritaria</i>	TTAGG	SART - (TTAGG)n - TRAS	OU538788.1-OU538819.1
Geometridae	<i>Crocallis elinguarua</i>	TTAGG	SART - (TTAGG)n - TRAS	OU026065.1-OU026082.1
Geometridae	<i>Ennomos fuscantarius</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992023.1-HG992054.1
Geometridae	<i>Ennomos quercinarius</i>	TTAGG	SART - (TTAGG)n - TRAS	OU342488.1-OU342519.1
Geometridae	<i>Erannis defoliaria</i>	TTAGG	SART - (TTAGG)n - TRAS	FR990066.1-FR990095.1
Geometridae	<i>Hydriomena furcata</i>	TTAGG	SART - (TTAGG)n - TRAS	OU538820.1-OU538848.1
Geometridae	<i>Hylaea fasciaria</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990192.1-LR990223.1
Geometridae	<i>Idaea aversata</i>	TTAGG	SART - (TTAGG)n - TRAS	OU026083.1-OU026113.1
Geometridae	<i>Peribatodes rhomboidaria</i>	TTAGG	SART - (TTAGG)n - TRAS	OU452166.1-OU452197.1
Hesperiidae	<i>Erynnis tages</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990071.1-LR990102.1
Hesperiidae	<i>Hesperia comma</i>	TTAGG	(TTAGG)n - TRAS	FR990012.1-FR990041.1
Hesperiidae	<i>Ochlodes sylvanus</i>	TTAGG	SART - (TTAGG)n - TRAS	FR990122.1-FR990152.1
Hesperiidae	<i>Pyrgus malvae</i>	TTAGG	SART - (TTAGG)n - TRAS	OU426946.1-OU426977.1
Hesperiidae	<i>Thymelicus sylvestris</i>	TTAGG	SART - (TTAGG)n - TRAS	OU426885.1-OU426913.1
Lycaenidae	<i>Aricia agestis</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990256.1-LR990279.1
Lycaenidae	<i>Celastrina argiolus</i>	TTAGG	SART - (TTAGG)n - TRAS	LR994577.1-LR994603.1
Lycaenidae	<i>Cyaniris semiargus</i>	TTAGG	SART - (TTAGG)n - TRAS	LR994546.1-LR994570.1
Lycaenidae	<i>Glaucopsyche alexis</i>	TTAGG	SART - (TTAGG)n - TRAS	FR990042.1-FR990065.1
Lycaenidae	<i>Lycaena phlaeas</i>	TTAGG	SART - (TTAGG)n - TRAS	HG995163.1-HG995187.1
Lycaenidae	<i>Lysandra bellargus</i>	TTAGG	SART - (TTAGG)n - TRAS	HG995319.1-HG995365.1
Lycaenidae	<i>Lysandra coridon</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992055.1-HG992145.1
Lycaenidae	<i>Plebejus argus</i>	TTAGG	SART - (TTAGG)n - TRAS	FR989926.1-FR989949.1
Noctuidae	<i>Abrostola tripartita</i>	TTAGG	SART - (TTAGG)n - TRAS	HG996486.1-HG996517.1
Noctuidae	<i>Acronicta aceris</i>	TTAGG	SART - (TTAGG)n - TRAS	OU342756.1-OU342788.1
Noctuidae	<i>Agrochola circellaris</i>	TTAGG	SART - (TTAGG)n - TRAS	OU611839.1-OU611869.1
Noctuidae	<i>Amphipyra berbera</i>	TTAGG	SART - (TTAGG)n - TRAS	OU343121.1-OU343152.1
Noctuidae	<i>Amphipyra tragopoginis</i>	TTAGG	SART - (TTAGG)n - TRAS	HG991991.1-HG992022.1
Noctuidae	<i>Apamea monoglypha</i>	TTAGG	SART - (TTAGG)n - TRAS	OU426914.1-OU426945.1
Noctuidae	<i>Atethmia centrago</i>	TTAGG	SART - (TTAGG)n - TRAS	HG995366.1-HG995396.1
Noctuidae	<i>Autographa gamma</i>	TTAGG	SART - (TTAGG)n - TRAS	LR989849.1-LR989881.1
Noctuidae	<i>Autographa pulchrina</i>	TTAGG	SART - (TTAGG)n - TRAS	FR997761.1-FR997793.1
Noctuidae	<i>Cosmia trapezina</i>	TTAGG	SART - (TTAGG)n - TRAS	LR991019.1-LR991051.1
Noctuidae	<i>Craniophora ligustri</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990954.1-LR990986.1
Noctuidae	<i>Dryobotodes eremita</i>	TTAGG	SART - (TTAGG)n - TRAS	OU823241.1-OU823273.1

Noctuidae	<i>Eupsilia transversa</i>	TTAGG	SART - (TTAGG)n - TRAS	OU611871.1-OU611903.1
Noctuidae	<i>Griposia aprilina</i>	TTAGG	SART - (TTAGG)n - TRAS	OU744283.1-OU744315.1
Noctuidae	<i>Hecatera dysodea</i>	TTAGG	SART - (TTAGG)n - TRAS	HG995286.1-HG995318.1
Noctuidae	<i>Hydraecia micacea</i>	TTAGG	SART - (TTAGG)n - TRAS	OU611774.1-OU611806.1
Noctuidae	<i>Mamestra brassicae</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990987.1-LR991018.1
Noctuidae	<i>Mesoligia furuncula</i>	TTAGG	SART - (TTAGG)n - TRAS	OU744790.1-OU744821.1
Noctuidae	<i>Mythimna ferrago</i>	TTAGG	SART - (TTAGG)n - TRAS	OU342673.1-OU342705.1
Noctuidae	<i>Mythimna impura</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990339.1-LR990371.1
Noctuidae	<i>Noctua fimbriata</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990921.1-LR990953.1
Noctuidae	<i>Noctua janthe</i>	TTAGG	SART - (TTAGG)n - TRAS	OU342552.1-OU342583.1
Noctuidae	<i>Noctua pronuba</i>	TTAGG	SART - (TTAGG)n - TRAS	LR999891.1-LR999923.1
Noctuidae	<i>Ochropleura plecta</i>	TTAGG	SART - (TTAGG)n - TRAS	FR997721.1-FR997753.1
Noctuidae	<i>Omphaloscelis lunosa</i>	TTAGG	SART - (TTAGG)n - TRAS	OU744243.1-OU744275.1
Noctuidae	<i>Phlogophora meticulosa</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990517.1-LR990548.1
Noctuidae	<i>Xestia c-nigrum</i>	TTAGG	SART - (TTAGG)n - TRAS	OU745243.1-OU745274.1
Noctuidae	<i>Xestia xanthographa</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990641.1-LR990672.1
Notodontidae	<i>Clostera curtula</i>	TTAGG	SART - (TTAGG)n - TRAS	FR997794.1-FR997824.1
Notodontidae	<i>Furcula furcula</i>	TTAGG	SART - (TTAGG)n - TRAS	OU452242.1-OU452271.1
Notodontidae	<i>Notodonta dromedarius</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990159.1-LR990190.1
Notodontidae	<i>Phalera bucephala</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990609.1-LR990640.1
Notodontidae	<i>Pheosia gnoma</i>	TTAGG	SART - (TTAGG)n - TRAS	FR989894.1-FR989925.1
Notodontidae	<i>Pheosia tremula</i>	TTAGG	SART - (TTAGG)n - TRAS	HG995397.1-HG995428.1
Notodontidae	<i>Ptilodon capucinus</i>	TTAGG	SART - (TTAGG)n - TRAS	OU611807.1-OU611838.1
Nymphalidae	<i>Boloria selene</i>	TTAGG	SART - (TTAGG)n - TRAS	HG993131.1-HG993161.1
Nymphalidae	<i>Erebia ligea</i>	TTAGG	SART - (TTAGG)n - TRAS	OU785219.1-OU785248.1
Nymphalidae	<i>Fabriciana adippe</i>	TTAGG	SART - (TTAGG)n - TRAS	FR989982.1-FR990011.1
Nymphalidae	<i>Limenitis camilla</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990224.1-LR990255.1
Nymphalidae	<i>Maniola jurtina</i>	TTAGG	(TTAGG)n - TRAS	HG995207.1-HG995237.1
Nymphalidae	<i>Melitaea cinxia</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992209.1-HG992240.1
Nymphalidae	<i>Mellicta athalia</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992176.1-HG992208.1
Nymphalidae	<i>Nymphalis io</i>	TTAGG	SART - (TTAGG)n	LR989895.1-LR989926.1
Nymphalidae	<i>Nymphalis polychloros</i>	TTAGG	SART - (TTAGG)n - TRAS	HG992241.1-HG992273.1
Nymphalidae	<i>Nymphalis urticae</i>	TTAGG	SART - (TTAGG)n - TRAS	LR989982.1-LR990014.1
Nymphalidae	<i>Pararge aegeria</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990891.1-LR990920.1
Nymphalidae	<i>Vanessa atalanta</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990549.1-LR990581.1
Nymphalidae	<i>Vanessa cardui</i>	TTAGG	SART - (TTAGG)n - TRAS	LR999924.1-LR999956.1
Peleopodidae	<i>Carcina quercana</i>	TTAGG	SART - (TTAGG)n - TRAS	OU342426.1-OU342456.1
Pieridae	<i>Anthocharis cardamines</i>	TTAGG	SART - (TTAGG)n - TRAS	FR989950.1-FR989981.1
Pieridae	<i>Aporia crataegi</i>	TTAGG	SART - (TTAGG)n - TRAS	OU538729.1-OU538755.1
Pieridae	<i>Colias croceus</i>	TTAGG	SART - (TTAGG)n - TRAS	HG991958.1-HG991990.1
Pieridae	<i>Pieris brassicae</i>	TTAGG	SART - (TTAGG)n - TRAS	LR989932.1-LR989948.1
Pieridae	<i>Pieris napi</i>	TTAGG	SART - (TTAGG)n - TRAS	HG993162.1-HG993187.1
Pterophoridae	<i>Emmelina monodactyla</i>	TTAGG	SART - (TTAGG)n - TRAS	OU745296.1-OU745326.1
Pyrilidae	<i>Endotricha flammealis</i>	TTAGG	SART - (TTAGG)n - TRAS	LR990852.1-LR990884.1

Sesiidae	<i>Bembecia ichneumoniformis</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342520.1-OU342551.1	
Sesiidae	<i>Sesia apiformis</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU611947.1-OU611978.1	
Sesiidae	<i>Synanthedon vespiformis</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU906945.1-OU906976.1	
Sphingidae	<i>Deilephila porcellus</i>	TTAGG	SART - (TTAGG) _n - TRAS	LR999970.1-LR999999.1	
Sphingidae	<i>Hemaris fuciformis</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU015431.1-OU015461.1	
Sphingidae	<i>Laothoe populi</i>	TTAGG	SART - (TTAGG) _n - TRAS	HG992146.1-HG992175.1	
Sphingidae	<i>Mimas tiliae</i>	TTAGG	SART - (TTAGG) _n - TRAS	HG995238.1-HG995267.1	
Tineidae	<i>Tinea semifulvella</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342584.1-OU342629.1	
Tineidae	<i>Tinea trinotella</i>	TTAGG	SART - (TTAGG) _n - TRAS	HG992305.1-HG992335.1	
Tortricidae	<i>Apotomis turbidana</i>	TTAGG	SART - (TTAGG) _n - TRAS	LR990280.1-LR990308.1	
Tortricidae	<i>Cydia splendana</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342871.1-OU342899.1	
Tortricidae	<i>Hedya salicella</i>	TTAGG	SART - (TTAGG) _n - TRAS	FR990096.1-FR990121.1	
Tortricidae	<i>Notocelia uddmanniana</i>	TTAGG	SART - (TTAGG) _n - TRAS	LR991052.1-LR991080.1	
Tortricidae	<i>Pammene fasciana</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU452272.1-OU452300.1	
Ypsolophidae	<i>Ypsolopha scabrella</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU342960.1-OU342991.1	
Zygaenidae	<i>Zygaena filipendulae</i>	TTAGG	SART - (TTAGG) _n - TRAS	OU015649.1-OU015680.1	
TRICHOPTERA					
Limnephilidae	<i>Limnephilus lunatus</i>	TTAGG	Various sequences with predominance of 166 bp and 22 bp repeats or (TTAGG) _n – TRAS structure (TTAGG) _n – TRAS	OU830592.1-OU830604.1	
Limnephilidae	<i>Limnephilus rhombicus</i>	TTAGG		OV815270.1- OV815276.1	
Limnephilidae	<i>Limnephilus marmoratus</i>	TTAGG		(TTAGG) _n – TRAS	OU862906.1-OU862920.1
COLEOPTERA					
Attelabidae	<i>Apoderus coryli</i>	TTGGG	10 bp short variable subtelomeric repeats (TTGACTCAC, TTTTACTCAC, TTTGACTCAT, TTGACTCTC, or similar); or short variable subtelomeric repeats TTTAATTA (and similar); or long repeats (chromosome 6, right), or different repeats, or SART (chromosome 6 left). In the chromosomes 6, 7, 11, 12, 14, 16, 18, subtelomeric sequences are found at the very ends of the chromosomes (as telomeric elements)	OU452198.1-OU452217.1	
Cantharidae	<i>Cantharis rustica</i>	TTAGG		SART - (TTAGG) _n	OU426877.1-OU426883.1
Cantharidae	<i>Rhagonycha fulva</i>	TTAGG		SART - (TTAGG) _n	HG996554.1-HG996560.1
Carabidae	<i>Leistus spinibarbis</i>	TTAGG		SART - (TTAGG) _n	OW121791.1-OW121792.1
Carabidae	<i>Pterostichus madidus</i>	TTAGG		SART - (TTAGG) _n	OU452301.1-OU452319.1
Coccinellidae	<i>Adalia bipunctata</i>	TTAGG		SART - (TTAGG) _n	OU342948.1-OU342958.1
Coccinellidae	<i>Coccinella septempunctata</i>	TTAGG		SART - (TTAGG) _n (found only in the right telomere of the chromosome 8)	OU015573.1-OU015581.1
Coccinellidae	<i>Harmonia axyridis</i>	TTAGG		SART - (TTAGG) _n	OU611927.1-OU611934.1
Elateridae	<i>Agrypnus murinus</i>	TTAGG			OV816024.1- OV816023.1
Melyridae	<i>Malachius bipustulatus</i>	TCAGG		Various repeats	OU342630.1-OU342639.1
Pyrochroidae	<i>Pyrochroa serraticornis</i>	TCAGG	SART - (TCAGG) _n	HG995152.1-HG995161.1	
Staphylinidae	<i>Ocypus olens</i>	TTAGG	SART - (TTAGG) _n	OU343047.1-OU343065.1	
Staphylinidae	<i>Philonthus cognatus</i>	TTAGG interspersed with TTTAGG	SART - (TTAGG)	OW052241.1-OW052250.1	

HYMENOPTERA				
Andrenidae	<i>Andrena dorsata</i>	TTAGTTTGGG		OV815487.1- OV815489.1
Andrenidae	<i>Andrena haemorrhoa</i>	TTAGGTCTGGG	(TTAGGTCTGGG) _n - combination of different short and medium-sized repeats - (TTAGGTCTGGG) _n , TTAGGTCTGGG - SART - TTAGGTCTGGG, combination of different short and medium-sized repeats	OU342940.1-OU342946.1
Andrenidae	<i>Andrena minutula</i>	TTAGGG	Various short and medium repeats	OV815993.1- OV815999.1
Apidae	<i>Apis mellifera</i>	The main repeat TTAGG interspersed with variant repeats TTAGGGTT, TTATAGG	15 bp repeat (TTAGGTCAGGCTGGG) _n , 20 bp repeat (TTAGGCTAGGTCAGGCTGGG) _n (chromosome 1, left telomere)	CM009931.2 - CM009946.2
Apidae	<i>Bombus campestris</i>	The telomere consists of the main repeat TTAGGTTGGGG interspersed with variant repeats TTAGGTGGGGG, TTAGGTTGGTGGG G, TTAGGTTGGTTAG TGGGG, TTAGGTTGGGTAG GTTGG, TTAGGTTGGGT, TTTGGGTGGGG TTAGGTTGGGG	SART - (TTAGGTTGGGG) _n	HG995126.1-HG995150.1
Apidae	<i>Bombus hortorum</i>	TTAGGTTGGGG	SART - (TTAGGTTGGGG) _n	HG995188.1-HG995205.1
Apidae	<i>Bombus hypnorum</i>	TTAGGTTGGGG	SART - (TTAGGTTGGGG) _n	OU427020.1-OU427031.1
Apidae	<i>Bombus pascuorum</i>	TTAGGTTGGGG	SART - (TTAGGTTGGGG) _n	HG995268.1-HG995284.1
Apidae	<i>Bombus pratorum</i>	TTAGGTTGGGG	SART - (TTAGGTTGGGG) _n	OV883983.1-OV884000.1
Apidae	<i>Bombus sylvestris</i>	TTAGGTTCCGGG (majority) interspersed with TTAGGTTAGGTTCCGGG	Various repeats with significant inclusion of the 41 bp motif TTCCAAATCTGTTTCAGATCCATCCCGGTTTCAGTCCAGACCG (and its variations)	OU443141.1-OU443164.1
Apidae	<i>Nomada fabriciana</i>	TTAGG	Various short, medium and long repeats	OU015688.1-OU015699.1
Bembicidae	<i>Nysson spinosus</i>	TTAGGTCTGGG	Combination of various short and medium repeats	OU342830.1-OU342855.1
Crabronidae	<i>Ectemnius continuus</i>	TTAGGTCTGGG	Combination of various short and medium repeats	OU342856.1-OU342869.1
Crabronidae	<i>Ectemnius lituratus</i>	TTAGGTCTGGG	TTAGGTGTGGG, TTTTCCTGGAA, TTTTCCTGGAATTTCCCGCGG (and other variations)	OU343033.1-OU343045.1
Formicidae	<i>Formica selysi</i>	TTAGG	TTAGCACGGGGTTACGG, TTAGGGTTAGCATGGGG, TTTGGTCAGGCTAGGCAAGGCAATG TCAGA (and other variations)	CM020805.1-CM020831.1
Formicidae	<i>Solenopsis invicta</i>	TTAGG	TTTGGTTATGGTTT, TTTGGTTATGGTCC, TTTGGTTATGGTCT (and other variations)	CM028732.1-CM028747.1
Halictidae	<i>Lasioglossum lativentre</i>	TTAGGTCTGGG (see Table 1)	see Table 1	OU744355.1-OU744368.1
Halictidae	<i>Lasioglossum morio</i>	TTAGG	22 bp motif TTAGGTTAGCGGATTCCGGG (variant TTAGGTTAGAGCGATTTCCGAG in the chromosome 1); however, the chromosome 7 has another subtelomere organization: combination of various short and medium motifs just before telomere motif (TTAGG) _n	OU744323.1-OU744334.1
Halictidae	<i>Lasioglossum pauxillum</i>	TTAGG	SART - (TTAGG) _n	OW121699.1- OW121707.1
Halictidae	<i>Seladonia tumulorum</i>	TTAGGG	Combination of various short, medium and long repeats with prevalence of TTTTTCGCGTCCGTAGAAGG, TTTTTCGAGGTCGGGGTGGT,	OU565266.1-OU565282.1

Halictidae	<i>Sphcodes monilicornis</i>	TTAGGTCTGGG	TTTTTGCGTCCGTAGAAGGTTG (and similar) TTAGGCCCTGCTCGG, TTAGGCCTGCTCGGT	OU565284.1-OU565302.1
Ichneumonidae	<i>Amblyteles armatorius</i>	T	18 bp subtelomeric repeat CAGCGCCACAGCGTCCG and similar repeats	OW121686.1- OW121697.1
Ichneumonidae	<i>Buathra laborator</i>	TTCCTC	Various sequences	OW203892.1- OW203902.1
Ichneumonidae	<i>Ichneumon xanthorius</i>	TTAAAACGCC, TTGAAACGCC, TTAAAGCGCC, TGGATGGGA, TTAAGCGTT, AAAAAATTCTTTG ATGC (see also text)	Various short, medium and long repeats	OU824200.1-OU824211.1
Melittidae	<i>Macropis europaea</i>	The motives TTGGG and TTAGG are mixed. The TTGGG motif prevails. The TTAGG motif almost never occurs twice in a row. TTGGG motif up to 13 times in a row.	121bp repeat	OU744343.1-OU744353.1
Pemphredonidae	<i>Mimumesa dahlboni</i>	TTAGG	Combination of various short and medium repeats; SART - (TTAGG) _n (chromosome 9)	OU824113.1-OU824130.1
Pemphredonidae	<i>Pemphredon lugubris</i>	TCTGGG	SART - (TTAGG) _n	OW121815.1-OW121819.1
Philanthidae	<i>Cerceris rybyensis</i>	TTAGGTCTGGG	TTAGGTGGTTGGACATG, TTAGGTGGGTGGACATG, TTAGGTGGTTGGACATG	OU342789.1-OU342802.1
Pompilidae	<i>Anoplius nigerrimus</i>	TTAGGTCTGGG	Combination of various short, medium and long repeats	OU612077.1-OU612091.1
Pteromalidae	<i>Nasonia vitripennis</i>	Within the main motif TTATTGGG , which obviously prevails, modified sequences are regularly found - TTATTGGGGG, TTTTTATTGGG, TTTTTATTGGG, TTATTGG, TTATTGGAGGG, TTATTGGGG, TTATTATTGGG, TTTTGTTATTGGG TTAGG	Combination of various short, medium and long repeats	CM020934.1-CM020938.1
Tenthredinidae	<i>Tenthredo notha</i>	TTAGG	Various short, medium and long repeats; TTAGG interspersed by various repeats; SART - (TTAGG) _n	OU611906.1-OU611925.1
Vespidae	<i>Ancistrocerus nigricornis</i>	TTGGGTCTGGG	SART - (TTGGGTCTGGG) _n	OU696664.1-OU696669.1
Vespidae	<i>Dolichovespula saxonica</i>	TTGCGTCTGGG	31 bp motif: TTTTGGACACTACCGCCTTTCGTA ACAGTA, TTTAGACTACTACGCCTTCATAA GAGTAG, TTTTGGACACTACCGCCTTTCGTA ACAGTA, TTTTAGAGACTACCGCCTTTCGTA ACAGTA and other variants	OU426993.1-OU427018.1
Vespidae	<i>Dolichovespula media</i>	TTGCGTCTGGG	TTTTAGTTTACGCTCGCGTCTTGGGA, TTTTAGTTTACGCTCGCGTCTTGGGA, TTTTTGGTTTTCGGTCGCGTCTGGG, CCCCGCGTATGGGTTGCTCTG	OU426851.1-OU426875.1
Vespidae	<i>Dolichovespula sylvestris</i>	TTGCGTCTGGG	The telomeric repeat TTGCGTCTGGG, (TTGCGTCTGGG) ₂ (TTGCGTCTGGG) ₃ is often found in the subtelomeric region	OU964961.1- OU964986.1
Vespidae	<i>Vespa crabro</i>	TTGCGTCAGGG	TTTTTCTGGTTCCTTCCGGAGTTGT GACTGGTTACGTTTGTAG	OU342400.1-OU342424.1

Vespidae	<i>Vespa velutina</i>	TTGCGTCAGGG	AAAAGTTGTGTCTGGTTGCGACTGC AGTTGCGTCTGGTTACGTCCGGGTA GCGTCAGCGTAGCGACTGACGTTGC	OU525123.1-OU525147.1
Vespidae	<i>Vespula germanica</i>	TTGCGTCTGGG	GTCTGGTAGTGTTT TTGCGTCTGAAGG	HG996528.1-HG996552.1
Vespidae	<i>Vespula vulgaris</i>	TTGCGTCTGGG	TTGGGTCTGAGG, TTGGGTCTCAGG, TTGGGTCTGAAGG	FR997668.1-FR997692.1
DIPTERA				
Conopidae	<i>Sicus ferrugineus</i>	374 bp sequence		OV277348.1-OV277354.1
Sciomyzidae	<i>Coremacera marginata</i>	371 bp sequence		OU612043.1-OU612093.1
Syrphidae	<i>Melanostoma mellinum</i>	327-329 bp sequence		OU612058.1-OU612063.1
Syrphidae	<i>Scaeva pyrastris</i>	327 bp sequence		LR989927.1-LR989931.1
Tachinidae	<i>Tachina fera</i>	173-178 bp sequence		LR999963.1-LR999969.1