Heat shock regulatory elements are present in telomeric repeats of Chironomus thummi

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ABSTRACT

As in other Diptera, the telomeres of Chironomus thummi lack canonical short telomerase-specified repeats and instead contain complex sequences. They react to heat shock and other stress treatments by forming giant puffs at some chromosome termini, which are visible in polytene cells. All telomeres, except the telocentric end of chromosome four (4L), consist of large blocks of repeats, 176 bp in length. Three subfamilies of telomeric sequences have been found to show different distribution patterns between chromosome ends. TsA and TsC are characteristic of telomeres 3R and 4R, respectively, whereas TsB is present in the other non-telocentric telomeres. Heat shock transcription regulatory elements have been identified in the telomeric sequences, appearing differentially represented in the three subfamilies, but otherwise rather similar in size and sequence. Interestingly, TsA and TsB repeats share the well-conserved heat shock element (HSE) and GAGA motif, while the TATA box is only present in the former. Neither a HSE nor a TATA box appear in TsC repeats. Moreover, experimental data indicate that the HSE is functionally active in binding heat shock transcription factor (HSF). These results provide, for the first time, a molecular basis for the effect of heat shock on C. thummi telomeres and might also explain the different behaviour they show. A positive correlation between the presence of HSE and telomeric puffing and transcription under heat shock was demonstrated. This was also confirmed in the sibling species Chironomus piger. The significance of heat shock activation of telomeric repeats in relation to telomeric function is unknown at present, but it might be compared to the behaviour of other non-heat shock protein coding sequences, such as SINE-like and LINE-like retroelements, which have been reported to be activated by stress.

INTRODUCTION

In most organisms studied telomeric DNA consists of short (5–26 bp) tandem repeats maintained by telomerase, a special cellular reverse transcriptase responsible for RNA-templated regeneration of telomeres to counteract the loss of sequences due to incomplete replication at the DNA termini (reviewed in 1,2). However, there is increasing evidence for alternative telomere structures and different mechanisms for telomere maintenance. Telomeres in Drosophila melanogaster and its close relatives are elongated by retrotransposition of the terminus-specific HeT-A and TART non-LRT transposons (reviewed in 3). Telomeres with long complex tandem repeats have been found in other Diptera such as Chironomus (4,5), Anopheles (6) and, more recently, in Drosophila virilis and its sibling species (7). The same has also been found in some plants, for example members of the genera Allium and Aloe (8,9), and in the ciliate linear mitochondrial chromosome (10). These organisms may somehow represent an intermediate situation between short telomeric repeats and the long D. melanogaster retrotransposons, and deserve more detailed analysis. Chironomus telomeres offer an interesting model since different species have been extensively studied. In Chironomus pallidivitatus, 340 bp repeats reach the very end of the chromosome (11). Similar sequences, likely to be terminal, have been found in Chironomus thummi (176 bp), Chironomus tentans (350 bp) and Chironomus dilutus (341 bp) (4,12). Chironomid telomeres might regenerate DNA by a process different to that already characterised. In chironomids, telomeric repeats are transcribed (4,5) and large telomeric transcripts have been found in salivary glands and other larval tissues (13). Support for RNA-templated regeneration is found in the presence of reverse transcriptase-like activity in telomeric areas (14).

Chironomus thummi telomeres are particularly interesting since they may react to heat shock and other stress treatments by inducing terminal puffs. At telomere 3R, a prominent Balbiani ring-shaped puff named TBR is formed in polytene chromosomes. In cytological studies, TBRs show signs of transcriptional activation involving telomeric repeats. Upon induction, TBR binds specific factors such as Ku (15), Hsp90 (16), heat shock transcription factor (HSF) (13) and reverse transcriptase (14), an indication of the complex processes at this telomeric structure. However, five non-telocentric telomeres react weakly to heat shock. Finally, the 4R telomere and the 4L...
telocentromere seem to be refractory to heat shock. The significance of this particular behaviour of *C. thummi* telomeres, in which the heat shock response appears to be associated with telomeric organisation and/or function, is still to be elucidated. Recently, we described three different subfamilies of telomeric repeats (17). In the present investigation, the telomeric organisation of *C. thummi* was further characterised and a sequence showing high homology to the heat shock element (HSE) in some subfamilies of repeats was found. This element, highly represented in the telomeric blocks, was able to bind specifically to HSF in vitro. The existence of telomeric HSE, along with other regulatory elements such as TATA and GAGA boxes, may explain why *C. thummi* telomeres react to heat shock and other stress treatments.

**MATERIALS AND METHODS**

**Animals**

The experimental animals were fourth instar larvae of *C. thummi*, *Chironomus piger* and *C. thummi × C. piger* hybrids. These were raised under standard conditions at 18°C. Larvae were heat shocked at 35°C for 1 or 2 h in a preheated and aerated culture medium.

**Immunocytochemistry**

Salivary glands were isolated, fixed and stained with carmine:orcein as previously described (17). Squashes were fixed in ethanol:acetic acid (3:1) were used for DNA–RNA detection. These were rehydrated and denatured for 10 min at 40°C in 50% acetic acid. Slides were then blocked for 10 min at room temperature in phosphate-buffered saline (PBS), 0.1% Tween-20 (PT) and 2% low fat powdered milk and incubated for 1 h at room temperature with anti-DNA–RNA hybrid IgG antibodies. After two washes in PT they were then incubated with anti-goat IgGs conjugated with fluorescein isothiocyanate (FITC) (Sigma). Squashes fixed in ethanol:acetic acid (3:1) were denatured. Slides were rehydrated and incubated with 100 mg/ml of anti-HSF IgG (18) diluting in PT with 1% blocking reagent (Roche) for 1 h at room temperature. After washing in PT, the secondary antibody used was anti-rabbit IgG conjugated with FITC (Sigma). Slides were finally mounted in 9:1 glycerol: PBS with 1 mg/ml p-phenylene diamine and observed under a Zeiss photomicroscope equipped with a CCD camera. Images were captured using a IP-lab spectrum program and processed with Adobe PhotoShop 4.0.

**DNA extraction and Southern blotting**

Genomic DNA was extracted from 100 frozen larvae as described (17). Genomic and plasmid DNA digested with Rsal was electrophoresed in 1.5% agarose gels in 1× TBE (89 mM Tris, 89 mM boric acid, 2 mM EDTA) for 3 h at 60 V. Alkaline transfer onto positively charged nylon was performed following the manufacturer’s instructions (Boehringer). An Rsal fragment obtained by digestion of the TsA dimer was randomly primed with DIG High Prime (Boehringer), whereas the HSE probe was obtained by annealing oligonucleotides HSE29A (5′-CCCTCTAGGAA AATTCTAGA AAAAT CG-3′) and HSE29B (5′-CTCGATTTCTAGA ATTTTCCATA GG-3′). It was then labelled with 11-Dig-ddUTP by terminal transferase (Roche) following the manufacturer’s instructions. Hybridisation and detection was performed using CDP-Star (Boehringer) diluted 1:1000 in 1% diethanolamine.

**Electrophoretic mobility shift assay (EMSA)**

The probe for the EMSAs was prepared by annealing complementary synthetic oligonucleotides followed by end-labelling with [32P]ATP (Amersham) and polynucleotide kinase (Roche). *Chironomus thummi* protein extracts were obtained from 10 control or treated larvae and human extracts were obtained from HeLa cells infected with either adenovirus or recombinant adenovirus carrying human Hsf1d202–316 (19). Extracts were quantified using the Bradford method (Bio-Rad). The binding reaction was performed by preincubating either 10 mg *C. thummi* proteins or 2.5 mg human proteins with 2 mg poly(dI·dC) in CB buffer (0.4 mM EDTA, 80 mM Tris–HCl, pH 8, 20 mM MgCl2, 2 mM DTT, 2 mM CaCl2) on ice for 20 min. Two nanograms of end-labelled double-stranded probe (50 000–100 000 c.p.m.) were added to the reaction mixture and incubated for 30 min at 4°C. Samples were electrophoresed in 4% polyacrylamide gels in 0.5× TBE (89 mM Tris–borate and 2 mM EDTA), 10% glycerol buffer, 175 V, at room temperature for 2.5 h. In competition experiments, a 50-fold excess of unlabelled oligonucleotides was incubated with the reaction mixture for 15 min at 4°C before addition of the radiolabelled probe. Supershift assays were performed by incubating an antibody against HSF-1 (Stressgen) overnight at 4°C with the protein extracts before the radiolabelled probe was added to the reaction mixture. HSE29 (5′-CCCTCTAGGAA AATTCTAGA AAAAT CGAG-3′) and HSE29C (5′-CCCTCTAGGAA AATTCTAGA AAAAT CGAG-3′) were used as competitors.

**In situ hybridisation**

Salivary glands fixed in ethanol:acetic acid (3:1) and squashed in 50% acetic acid were denatured. Each slide was treated with 100 nmol telomeric HSE oligonucleotide (the same as that used in the Southern blot experiments) 3′-labelled by terminal transferase with 11-Dig-ddUTP (Roche) or TsA monomer labelled by random priming with 11-dUTP-biotin (Enzo) and incubated overnight at 37°C in hybridisation buffer (50% deionised formamide and 4× SSC, 0.4% SDS). Slides were washed in 2× SSC for 5 min and in PT. Detection of the probe was achieved with anti-digoxigenin IgGs conjugated with rhodamine (Roche) or FITC-conjugated EstrAvidin (Sigma). After washing, the slides were stained with Hoechst, mounted in 9:1 glycerol: PBS with 1 mg/ml p-phenylene diamine and observed with a Zeiss photomicroscope equipped with a CCD camera. Images were captured using the IP-lab spectrum program and processed with Adobe PhotoShop 4.0.

**RESULTS**

**Presence of a HSE in telomeric sequences**

Three subfamilies of repeats, TsA, TsB and TsC, were characterised previously (17) by sequence analysis of single repeats (monomers). In the present investigation, a head-to-tail dimer of TsA was isolated from a genomic minilibrary (4) and TsB dimers by PCR. A closer inspection of the TsA dimer sequence
led to the identification, in the junction of the two repeats, of the sequence gGAAaaTTCtaGAAa, which is highly homologous to the consensus sequence for the HSE, a transcriptional regulatory sequence shared by all heat shock coding genes. The HSE consensus sequence shows a repeating array of the 5 bp sequence nGAAn, where each module is inverted relative to the one immediately adjacent. The minimum size of an HSE is three modules (nGAAnnTTCnnGAAn) (20). The sequence containing the telomeric HSE shows great similarity to an HSE from the D.melanogaster hsp22 promoter (21). The interest of this finding in the context of telomeric activation by heat shock prompted a search for this element in dimers of the other types of repeats (TsB and TsC). Figure 1 shows the dimer sequence alignment. Homology with the HSE consensus sequence was observed in TsB, although less similarity was seen with TsC, due to the presence of three base substitutions.

A detailed analysis of the sequences revealed other regulatory elements, including a TATA box with an internal GC pair only found in TsA (Fig. 1) and also a GAGA motif running from position 46 to 52 in TsA which was present in all three subfamilies of repeats. The telomeric GAGA motif is homologous to the consensus sequence of the GAGA transcription factor, (GA·TC)n, which is found in the promoters of several Drosophila genes (22).

Abundance and distribution of HSE in telomeres

An attempt was made to ascertain the abundance of the identified HSE in telomeric DNA. In Southern blots, genomic DNA was digested with RsaI to maintain the HSE within one monomer. Identical samples of RsaI-digested DNA were electrophoresed and hybridised separately with either the whole telomeric repeat or an oligonucleotide containing the telomeric HSE (Fig. 2). The signals obtained with both probes were comparable, suggesting HSE to be highly represented in the telomeric blocks of repeats.

To determine the location of HSE at the level of individual telomeres, FISH experiments were performed on polytene chromosomes using an oligonucleotide covering the HSE as probe. As shown in Figure 3, a clear and distinct signal at all non-telocentric telomeres was obtained except for 4R. These results are in good agreement with the prediction made from the dimer sequence analysis, since TsC sequences (which lack
HSE) are specific for the 4R telomere (17). Further, the strong fluorescent signal confirmed the Southern blot data (Fig. 2), indicating that HSE is abundant in telomeres of C.thummi.

**In vitro binding of HSF by telomeric HSE**

To learn whether HSE binds HSF, EMSAs were performed with an oligonucleotide that runs from position 166 to 194 (Fig. 1). This includes three modules with homology to the HSE consensus and an additional 7 nt in each direction, forming a total size of 29 nt in all. Incubation of labelled HSE29 with extracts from control C.thummi larvae showed no shift in mobility (Fig. 4A, lane 1). When protein extracts from heat shocked larvae were used, a band was seen (Fig. 4A, lane 2) which disappeared in experiments using a 50-fold excess of cold oligonucleotide as competitor (Fig. 4A, lane 3). Nevertheless, a 50-fold excess of HSE29C, an equivalent oligonucleotide covering the same zone from TsC repeat, did not abolish HSE binding (Fig. 4A, lane 4). These results show that the HSE sequences identified in the dimers of TsA and TsB telomeric units are able to bind HSF-1 in vitro, suggesting that this element may be functional in vivo, while TsC units are unable to bind HSF under the same conditions.

**Involvement of HSE in heat shock-induced puffing**

Patterns of puffing, transcription and HSF binding in individual telomeres were analysed in polytene cells of C.thummi under heat shock conditions (Table 1). The results show a positive correlation between the presence of heat shock regulatory elements and the ability of specific telomeres to react to heat shock. Particularly evident was the case of telomere 3R, characterised by prominent puff formation (TBR III), HSF binding and transcriptional activation upon heat shock (Fig. 5e and f). TsA telomeric sequences containing HSE, GAGA and TATA boxes are specifically found in this telomere. In contrast, at telomere 4R, characterised by TsC sequences lacking HSE, neither puffing nor any other sign of activation was found (Fig. 5e and f). Other non-telocentric telomeres containing HSE and characterised by the presence of TsB sequences showed rather
irregular behaviour upon heat shock. While slight signs of transcriptional activation were detected at these telomeres, puff formation and HSF binding were only occasionally observed. The correlation between HSE and telomeric activation was further confirmed by analysing the behaviour of \textit{C.piger} telomeres under heat shock. In this species, a TBR was specifically induced by temperature at telomere 4R (23). A FISH experiment using HSE oligonucleotide as probe showed a clear positive signal at telomere 4R in \textit{C.piger} polytene chromosomes but not in \textit{C.thummi} telomeres when probed in \textit{C.thummi} \texttimes \textit{C.piger} hybrids (Fig. 5a–c). Further, HSF accumulated at the 4R \textit{C.piger} telomere during heat shock (Fig. 5d and e). Thus, the presence of HSE, along with the capability of binding HSF, seem to be the basic features connected to full expression of telomeric puffing upon heat shock.

### DISCUSSION
The existence of heat shock regulatory elements in the telomeric sequences reported in this paper provides, for the first time, a molecular basis for understanding the singular behaviour of \textit{C.thummi} telomeres. These telomeres can be induced, by heat shock, to form giant, actively transcribing terminal puffs, named TBRs, which are most pronounced at the 3R chromosome end (24). Other chromosomal ends are only occasionally induced to puff, although signs of transcription could be detected under heat shock conditions. Finally, neither of the ends of chromosome 4 was found to respond to heat shock. Telomeric HSE, found both in the TsA and TsB subfamilies of repeats, is highly homologous to the HSE consensus sequence established for heat shock protein coding genes. Telomeric HSE is able to bind HSF \textit{in vitro}, as demonstrated by EMSA, and might be active \textit{in vivo}, as suggested by HSF immunolocalisation in heat shock-induced telomeres.

The presence of HSE appears to be a necessary condition for telomeric activation by heat shock. The lack of response at the 4R end correlates with the lack of signal in FISH when probed with HSE oligonucleotides consistent with the TsC content of this telomere (17). Furthermore, in the related species \textit{C.piger}, where telomere 4R always forms a puff after heat shock, HSE is also present and HSF localises at this site.

| Table 1. Telomeric activation at different chromosome ends in \textit{C.thummi} after 2 h of 35°C heat shock |
|---|---|---|---|---|---|---|
| Size of puff | ++ | ++ | – | – | – | ++ |
| Puffing frequency | M | M | L | L | H | N |
| Transcriptional activity | ++ | ++ | ++ | ++ | + | +++ |
| HSF association | – | + | – | – | – | + |

Puff size was measured as large (+++), medium (++), small (+) or not visible (–). Puffing frequency was considered high (H) when it reached >30%, medium (M) when between 10 and 30% and low when <10% (L). In telomere 4R no puffing was observed. Transcriptional activity was studied using an antibody against DNA–RNA hybrids. The frequency of telomeres with a positive signal (+, >50%; ++, >60%; ++++, >80%) is indicated. HSF association indicates the presence (+) or absence (–) of a positive signal. At least five different larvae and 30–40 chromosomes per preparation were scored in each case.
Other transcriptional regulatory elements such as TATA and GAGA motifs were also found in the telomeric sequences. They are differentially represented in TsA and TsB telomeric subfamilies of repeats, a possible explanation for the differences in heat shock activation observed among individual chromosome termini. Telomere 3R, which is always induced after heat shock, consists of large homogeneous blocks of tandem repeats of TsA sequences containing the minimal heat shock promoter elements (HSE, TATA and GAGA) capable of reacting to thermal stimuli (25). The lack of a TATA box in TsB sequences might be related to the more irregular reaction to heat shock of this group of telomeres. Other possibilities, such as chromatin structure constraints, cannot, however, be ruled out.

The alternative, that telomeric induction in C. thummi may be due to heat shock protein coding genes located at the chromosomeal termini, is remote in view of the present results. The existence of heat shock regulatory elements integrated in the telomeric structure, possibly acting as internal promoters, seems more likely. In fact, RNA complementary to the telomeric repeats has been detected both locally and in northern blots after heat shock induction (26).

Heat shock activation of telomeres has only been observed in C. thummi and its sibling species C. piger, and not in several other Chironomus species investigated to date. All are similarly organised as large blocks of complex AT-rich sequences, but they show differences in the size of repeats as well as strong divergence at the sequence level (27). Nevertheless, constitutive transcription of telomeric repeats appears as a common feature (4,5). It is tempting to speculate that transcription of telomeric sequences might be directly involved in telomere maintenance through an RNA-templated mechanism, as suggested by the presence of retrotranscriptase detected in situ at telomeres (14).

The fact that heat shock responsiveness seems to be restricted to some telomeres in C. thummi and its sibling species C. piger make doubtful its requirement for essential telomeric maintenance functions. Nevertheless, it is remarkable that heat shock also activates transcription of the Zepp element of Chlorella telomeres (28). Zepp is a retrotransposon-like LINE-like element that accumulates at Chlorella subtelomeric regions, forming the actual terminus on chromosome I, which lacks the telomeric short repeats (29).

A number of mobile and/or interspersed repetitive elements are activated by heat shock, such as certain SINEs (30–33) and retrotransposons (34,35). The activation of such elements has, until now, not been considered part of the cellular stress response. However, it has recently been suggested that transcriptional heat shock activation of SINEs could play a protective role (36). To our knowledge, C. thummi telomeric repeats are the first short repetitive DNA units in which heat shock regulatory elements have been identified. The physiological importance of telomeric heat shock activation remains to be determined.

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