Reverse transcriptase activity associated with haemic neoplasia in the soft-shell clam *Mya arenaria*

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ABSTRACT: Reverse transcriptase (RT) activity has been reported in bivalves affected by haemic neoplasia (HN). Since all retroviruses have RT, detection of RT activity was regarded as evidence for the retroviral etiology of HN. This study investigates the relationship between RT levels and the progress of HN as indicated by percentages of tetraploid cells in soft-shell clams *Mya arenaria*. The percentages of tetraploid cells were estimated by flow cytometry, and the RT levels were quantified using TaqMan product-enhanced RT (TM-PERT) assay. Results demonstrated that the amount of RT was positively correlated with the percentage of tetraploid cells circulating in clam haemolymph ($R^2 = 0.974, p < 0.001$). Compared to HN-negative clams (<5% tetraploid cells), 2 stages with significantly elevated levels of RT activity were observed: the first stage at ~10 to ~20% tetraploid cells, and the second at ~30 to ~80% tetraploid cells ($p < 0.01$). These data support the well-established fact from mammalian models that transformed cells express high levels of non-telomeric RT. The observed increase in RT levels at ~30% tetraploidy coincides with previously reported p53 gene expression. Taken together, this could indicate that using RT levels as an indicator of HN, ≥30% tetraploidy is the stage at which the disease process undergoes a change, and perhaps becomes irreversible.

KEY WORDS: *Mya arenaria* · Haemic neoplasia · Reverse transcriptase · Polyploidy · TaqMan · Product-enhanced reverse transcriptase assay · PERT

INTRODUCTION

Haemic neoplasia (HN) is a leukemia-like disease of the soft-shell clam *Mya arenaria*. The condition is often fatal and progressive in most of the locations where the disease has been reported (Barber 2004). In Prince Edward Island (PEI), Canada, massive mortalities of soft-shell clams were linked to high prevalence of HN (McGladdery et al. 2001). Different diagnostic techniques for the detection of HN based on the morphological distinction between normal and neoplastic haemocytes have been employed (Elston et al. 1992, McGladdery et al. 2001, Barber 2004). Neoplastic cells have large pleomorphic nuclei containing one or more nucleoli, with higher nucleo-cytoplasmic ratios. Given the change in ploidy of circulating haemocytes in diseased individuals, DNA content measurement using flow cytometry (FCM) has increasingly become a powerful tool for diagnosis of HN in bivalves (Harper et al. 1994, Reno et al. 1994, da Silva et al. 2005, Delaporte et al. 2008). The ploidy status in HN has been shown to vary among bivalve species. In *Mya arenaria*, HN
affected individuals have been found to show tetraploid cells with 1.25 to 2.05 times more DNA than normal diploid haemocytes (Reno et al. 1994).

Since the initial description of the disease (Farley 1969), its cause has not been clearly defined (Barber 2004). Viral infection, genetic profile, environmental changes and anthropogenic pollution have been proposed as the causative factors (Elston et al. 1992, McGladdery et al. 2001, Barber 2004). Retroviral etiology of HN has been the detection of reverse transcriptase (RT) activity in individuals (Romalde et al. 2007). The main evidence supporting a retroviral etiology of HN has been the detection of reverse transcriptase (RT) activity in individuals affected by this pathological condition in soft-shell clams (Medina et al. 1993, House et al. 1998) and in cockles, Cerastoderma edule (Romalde et al. 2007). Reverse transcriptase activity, however, is not exclusive to retroviruses. Other viruses (hepadnaviruses), some bacteria (e.g. strains of Myxobacteria and Escherichia coli), and also telomerases (a set of enzymes that lengthen the chromosome end (telomere) by reverse transcription of its own RNA; Flint et al. 2004) have RT activity. Moreover, RT-related sequences are found in cellular genomes generally known as retroelements (Flint et al. 2004). Expression of these endogenous cellular RT-coding genes is repressed in normal terminally differentiated cells, but can be active in tissues with rapidly dividing cells such as germ cells, embryos, and tumors (Spadafora 2004).

The main contention of this study is that the absence of conclusive evidence of an exogenous retrovirus in the presence of RT activity likely means that the RT detected in HN is expressed from sources other than exogenous retrovirus. Thus, quantification of RT activity at different stages of HN development may provide valuable information on possible sources of RT activity. Product-enhanced RT (PERT) assay has been used as a method for general detection of RT activity of both known and unknown retroviruses, which was subsequently modified as TaqMan PERT (TM-PERT). TM-PERT is a quantitative assay in which the resulting cDNA from RT activity is amplified and quantified by fluorogenic 5′-nuclease TaqMan PCR (André et al. 2000, Maudru & Peden 1998). The use of TM-PERT for quantitative detection of RT activity can help to distinguish the background RT signals arising from nonRT sources, such as some thermostable DNA polymerases released from lysed cells. Also, unlike conventional PERT, TM-PERT detects manganese- and magnesium-dependent RT with approximately equal sensitivity (Maudru & Peden 1998). Finally, the TM-PERT method allows quantification of RT activity over a wider range of activities than PERT does (Maudru & Peden 1998).

This study describes the relationship between RT levels and the progress of HN as indicated by the percentage of tetraploidy in circulating haemocytes from soft-shell clams.

MATERIALS AND METHODS

Samples. Two hundred clams were collected from North River (46° 15′ 01″ N, 63° 10′ 42″ W), Charlotte-town, PEI, Canada in October 2007. Following gradual acclimation from 5 to 18°C over 2 d, the clams were held in tanks at 18°C until use.

Flow cytometry. The FCM procedure was performed according to Delaporte et al. (2008). Briefly, haemolymph was withdrawn from the anterior adductor muscle of each clam using a 25-gauge needle fitted with a 3 ml syringe. A 0.5 ml haemolymph sample was fixed in 2.5 ml of 95% cold ethanol. After centrifugation at 400 × g for 10 min, cell pellets were resuspended and rehydrated in phosphate buffered saline (PBS, 0.01 M) for 30 min, followed by 2 washes in PBS (0.01M). The resuspended filtered cell pellets were treated with DNase-free RNase A (Sigma, R4875, 50 µg ml⁻¹) and stained with propidium iodide (PI, Sigma, P4170, 50 µg ml⁻¹). A specific FL2 detector (orange light, at a wavelength of 550 to 600 nm) of a FACSCalibur flow cytometer (BD BioSciences) was used to measure fluorescence of PI-stained cells. For each sample, 10 000 particles were counted at a low flow rate (15 µl min⁻¹). For each cell, a single electronic pulse of PI fluorescence was recorded. Each pulse was discriminated by its area, height and width. Based on these data, the tetraploid cells at the G2/M phase were distinguished from normal diploid cells (2N) at the G0/G1 phase, and from doublets of diploid cells (2 cells with the same DNA quantity (2N) stuck together) by plotting FL2-area vs. FL2-width on cytograms (Fig. 1a). To discriminate single cells from doublets, the R1 region was drawn on data scatter plots so that cell doublets were plotted to the right of R1 (Fig 1a). Also, PI fluorescence intensities of single cells were plotted on an FL2-area histogram in order to calculate the percentage of normal cells and tetraploid cells in the tested sample (Fig. 1b).

Quantification of RT activity. Samples: After individual assessment of all clams by FCM, clams were assigned to 1 of 7 groups based on the percentages of tetraploid cells in the haemolymph, for TM-PERT analysis (Table 1). Clams with <5% tetraploid cells were considered as HN-negative controls based on Delaporte et al. (2008). Three clams were selected for inclusion in each of the 6 subsequent groups, except the ~70% group, which contained only 2 clams. The detailed composition of the experimental groups and the percentages of tetraploid cells in clams are listed in...
Table 1. Clams containing between >5% and <20% tetraploid cells were presumed doubtful, and those with >20% tetraploid cells were presumed positive based on Delaporte et al. (2008).

**Preparation of samples:** From each selected individual, 1 ml of haemolymph was withdrawn from the anterior adductor muscle as described above. The haemolymph samples were centrifuged at 500 × g for 15 min, and supernatants were again centrifuged at 13400 × g for 5 min. Supernatants were then passed through 0.22 μm filters (VWR International), aliquoted, and stored at −80°C.

**TM-PERT assay:** Assay conditions, and primer and probe sequences, were adopted from Maudru & Peden (1998) with some modifications. The respective sequences of primers A and B were 5′-GCC TTA GCA GTG CCC TGT CT-3′ and 5′-AAC ATG CTC GAG GGC CTT A-3′, while that of the probe was FAM-5′-CCC GTG GGA TGC TCC TAC ATG TC-3′-BHQ1.

For reverse transcription, 0.4 μg of the bacteriophage MS2 genomic RNA template (Roche Applied Science) was mixed with 5 μM of primer A in RNase-free water, in a final volume of 2 μl per RT reaction. The mixture was heated at 85°C for 5 min, annealed at 37°C for 30 min, and kept at 4°C for 5 min. Then, 2 μl of clam haemolymph sample was added to the RT reaction mixture containing 50 mM Tris-HCl, 8 mM MgCl₂, 30 mM KCl, 1 mM dithioerythritol, 10 U RNAsin (Promega), and 1 mM dNTPs (Qiagen), to give a 20 μl final volume. The reaction was then incubated at 37°C for 1 h, then at 95°C for 7 min.

For PCR amplification, 5 μl of the synthesized cDNA was added to the PCR reaction mixture to give a total volume of 25 μl. The reaction mixture included 1× Taq-Man Universal Master Mix including AmpErase uracil-N-glycosylase (UNG) (Applied Biosystems), 0.3 μM of primer A, 0.3 μM of primer B, 0.15 μM of probe labeled at the 3′ end with FAM and at the 5′ end with Black Hole Quencher (BHQ) 1 (Biosearch Technologies), and 250 ng of RNase A (Qiagen). The tubes were placed in the Chromo 4 system (Bio-RAD), and incubations were controlled using MJ Opticon Monitor version 3.1 with the following thermal cycler conditions: 37°C for 15 min; 50°C for 2 min; 95°C for 10 min; 45 cycles at 95°C for 15 s, 56°C for 15 s, and 72°C for 30 s. A standard curve was established by using a 10-fold dilution of Moloney murine leukemia virus (M-MLV) RT (Roche Applied Science) from 10² to 10¹⁰ picounits (pU). The enzyme was diluted in buffer A consisting of 50 mM KCl, 20 mM Tris-HCl pH 7.5, 0.2 mM dithiothreitol, 0.25 mM EDTA, 0.025% Triton X-100 (v/v), and 50% glycerol (v/v). The amplified PCR products of the serial dilutions were resolved by 1.5% agarose gel electrophoresis in 1× tris-borate-EDTA (TBE) buffer, visualized by staining with ethidium bromide, and photographed using UV illumination.

**Electron microscopy.** Haemolymph samples from clams (n = 10) previously classified by FCM to be between 8 and 70% tetraploid, were analyzed by electron microscopy (EM) in order to explore the presence of retroviral particles. Haemolymph samples were fil-
tered using a 0.45 µm syringe filter (VWR International), and centrifuged at 100 000 × g for 90 min in an SW 60 Ti rotor (Beckman). Each resultant pellet was resuspended in 30 µl TNE buffer (10 mM Tris pH 7.4, 400 mM NaCl and 1 mM EDTA; Oprandy et al. 1981). Volumes of 10 µl were placed on Formvar-coated grids and allowed to dry. One drop of a mixture of phosphotungstic acid and bovine serum albumin was placed on the grid and the excess was blotted off with filter paper. The samples were examined with a Hitachi H7500 electron microscope (Nissei Sangyo) at 80 kV.

Data analysis. A linear regression line between the logarithms of the RT standards and the corresponding threshold cycle (C_T) values was calculated and plotted. The logarithms of RT levels in haemolymph samples were determined by extrapolating the C_T values from the standard curve. One-way ANOVA was performed to assess multiple comparisons of log RT concentrations among different tetraploidy groups using Minitab software version 15 (Minitab). STATA software version 9 (College Station, Texas, USA) was used for exponential regression analysis. The Gompertz model was used to assess correlation between % tetraploidy and log RT concentrations.

RESULTS

The FCM analysis showed that 116 (58%) clams were negative (<5% tetraploid cells), 42 (21%) were diseased (>20% tetraploid cells), 20 contained between 20 to 50% tetraploid cells, and 22 contained >50% tetraploid cells.

TM-PERT assay optimization enabled the detection of up to 10^2 pU of RT 2 µl^-1 of haemolymph or 5 × 10^9 pU ml^-1, which is equivalent to 50 retroviral particles ml^-1 (a retrovirus particle contains ~10^3 RT pU) (Maudru & Peden 1998). The M-MLV RT standard curve showed a linear relationship between threshold cycle and the log of RT concentrations over a wide range of concentrations (10^2 to 10^10 pU) (Fig. 2a). The numerical means and SD values of the slopes, the y-intercepts, and the R^2 of the assays performed were −3.631 ± 0.217, 46.64 ± 3.16, and 0.9965 ± 0.00214, respectively. The PCR product target specificity of TM-PERT assay was verified by agarose gel electrophoresis, which revealed a band of the expected size (112 bp). In addition, negative controls with no RT source, which were replaced with either RNase-free water or buffer A, showed no bands (Fig. 2b).

One of the HN-negative haemolymph samples used for quantitative assessment of RT activity exhibited no signal; the other 2 ind. showed C_T values of 34.31 and 39.58, respectively. The logarithms of RT levels in individuals with ~10 to ~80% tetraploid cells ranged from 3.6 to 6.6 pU 2 µl^-1 (Table 2), which are equivalent to 2 × 10^6 to 2 × 10^8 pU ml^-1 of haemolymph, respectively. Compared to the HN-negative group, a significant increase in RT levels was observed in clams with ~10 and ~20% tetraploidy, with no significant difference in RT levels between the 2 groups. Further significant increase in RT levels was observed in the haemolymph of clams with ~30% tetraploid cells and above (p < 0.01) (Table 2). Under the conditions of this study, clams could be classified into 3 distinct groups according to the RT levels: <5% tetraploid cells; ~10 to ~20% tetraploid cells; and ~30 to ~80% tetraploid cells. Overall, there was a significant positive correlation between the percentages of tetraploidy and RT levels (R^2 = 0.974, p < 0.001) (Fig. 3).
Electron microscopic examination of haemolymph samples using negative staining did not show retrovirus-like particles.

**DISCUSSION**

The FCM analysis of the clam population studied showed that 21% of tested clams were affected, which confirms previous reports by Delaporte et al. (2008) and Siah et al. (2008) suggesting that HN is widely spread among North River clams.

The PERT assay has been used as a method to quantify known and detect unknown retroviruses, with recognized limitations in terms of possible detection of RT-like activities of nonretroviral origin (Maudru & Peden 1997, André et al. 2000, Brorson et al. 2002). Apart from retroviruses, there are several other possible sources of RT activity. At the level of the assay itself, there are some reagents that may produce RT-like activity such as Taq DNA polymerases or RNase inhibitors (Pyra et al. 1994). At the level of the tested sample, there are some enzymes that may display RT-like activity such as host DNA polymerases, some other nuclear enzymes, and polymerase from mitochondria (Brorson et al. 2002).

In this study, in order to measure introduction of RT activity from reagents, negative controls with water and buffer A were always included. As a further control to measure possible contribution of sample-unrelated RT activity arising from PCR reagents, an RNA digestion step with RNase A was included before PCR amplification (Maudru & Peden 1997). In order to avoid false positive results possibly arising from released cellular enzymes, haemocytes were carefully removed from haemolymph samples by low speed centrifugation, and the assays were performed on a cell free haemolymph.

Although it could be argued that telomerase activity could be present since HN is widely believed to involve cell transformation, our assay should not have detected telomerase activity for 2 reasons. Firstly, telomerases are template specific, requiring a specific recognition sequence, TTAGGG. Thus, it is generally believed that they are unlikely to reverse transcribe MS2 phage RNA which lacks this target sequence (Brorson et al. 2002). Secondly, the assay was performed on cell free haemolymph as discussed above. On the other hand, TM-PERT is able to detect RT activity of endogenous retroviruses and other retroelements such as retrotransposons (Brorson et al. 2002).

Our detection of low levels of RT activity in some disease-negative clams is in agreement with findings in other species: e.g. in turtles (Casey et al. 1997), humans (Molès et al. 2007), and cockles (Romaule et al. 2007). The observed RT activity in HN-negative clams could be related to the fact that RT activity is associated with a wide range of biological processes, both physiological and pathological (Spadafora 2004). This is because RT-coding genes are also contained in repeated genomic elements called retroelements, which play an important role in many physiological and pathological cellular processes (Spadafora 2004).

We found a statistically significant difference in RT quantities between groups with ~10 to ~20% tetraploid cells and ~30 to ~80% tetraploid cells (p < 0.01) (Table 2). This supports previously published obser-
tions of disease remission in clams (Cooper et al. 1982, Leavitt et al. 1994) and mussels (Elston et al. 1988) in early stages of the disease (~20% neoplastic cells). The increased level of significance at ~30% tetraploidy suggests that this might be the beginning of the irreversible stage in the progression of the pathological process.

The absence of structures similar to retroviral particles in negatively stained samples of clam haemolymph is in disagreement with Oprandy et al. (1981) but in agreement with House et al. (1998) and other electron microscopic studies reviewed by Elston et al. (1992) and Barber (2004). The elevated levels of RT activity at various levels of tetraploidy without indication of retroviral particles in haemolymph samples as evaluated by electron microscopy suggests that the observed RT activity might be due to the expression of endogenous source of RT. Others demonstrated the presence of RT protein in tissues that were not included in retroviral particles, and they suggested endogenous retroelements as the source (Molés et al. 2007).

The positive correlation between levels of tetraploidy and RT activity suggests that the transformed haemocytes might have a role in the increased level of RT activity. Our suggestion is supported by the growing body of data showing that transformed cells of mammalian origin express high levels of endogenous non-telomeric RT, with the source of the RT being endogenous retroelements (Spadafora 2004, Oricchio et al. 2007). Involvement of endogenous retrovirus in HN of Mya arenaria was previously suggested by Oprandy & Chang (1983).

Further supporting our contention of an endogenous source of RT is the report that RT-coding genes are generally active in cancer cells (Spadafora 2004). In addition, RT gene activity is up-regulated by a variety of stimuli acting at the genomewide level such as cellular stress, heat shock, genotoxic agents and others (Sciamanna et al. 2005).

In conclusion, we suggest that RT activity associated with HN might be due to activation of endogenous retroelements, but whether this activity is a consequence, or is related to the cause of the disease requires further investigation. What retroelement may be responsible for production of RT activity is presently unclear. On the other hand, whether or not an exogenous retrovirus plays a role in induction of HN cannot be conclusively answered by our current data; however, our results on RT activity and EM analysis of haemolymph, combined with observations by other authors (Elston et al. 1992, House et al. 1998, Molés et al. 2007), suggest that the RT activity is not from an exogenous retrovirus.

Compared to HN-negative animals, the 2 significant increases in RT levels—first at ~10 to ~20% tetraploid cells and second at ~30 to ~80% tetraploid cells—suggest that the stage from ≤10 to ≥30% tetraploid cells should be targeted for further investigations on the mechanism of induction of HN.

Acknowledgements. We thank J. W. Casey for his helpful advice, G. Johnson for his valuable support of this study, and A. Mournchili for his help in statistical analysis. The research was supported by the Egyptian Cultural and Educational Bureau in Montréal, Quebec, Canada.

LITERATURE CITED

AboElkhair et al.: Reverse transcriptase activity in haemic neoplasia of *Mya arenaria*


Editorial responsibility: Catherine Collins, Aberdeen, UK

Submitted: August 4, 2008; Accepted: January 25, 2009
Proofs received from author(s): February 26, 2009