Correlation between Exons and Dispersed Repetitive DNA Distribution on the Human Genome

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1 Introduction

The human nuclear genome contains a large number of highly repeated DNA sequences. The Alu sequences are primate specific and are the most abundant family of repeated DNA sequences in the human genome. The human Alu sequence is approximate 300 bp long [2]. The L1 sequence is a long interspersed nuclear element. L1 is found in other mammals. Although their functions are not yet clear [4], some of them may affect gene functions or cause human diseases [3].

We have identified repeated DNA sequences from human genomic sequences in the region of 3p21.3p22 and 9q32, both of which are more than 1M bp long. Our statistical analysis shows that the distributions of *Alus* and exons have a weak positive correlation and those of *L1*s and exons have a weak negative correlation.

2 Method

Genomic sequence data of the human chromosome 3p21.3-p22 and 9q32 as well as cDNA sequences on these regions were obtained by Y.Daigo *et al.* (unpublished data). The lengths of sequences on the chromosome 3p21.3-p22 and 9q32 are 1.2M bp and 1.0M bp respectively. While the region 3p21.3-p22 contains 14 genes, the region 9q32 has only 3 genes.

Repetitive sequences were identified by the computer program CENSOR [1] with Repbase(Release 5.0). We divided each sequence into non-overlapping 100k bp segment and counted the exon, Alu and L1.

3 Result & Discussion

To characterize the exon, Alu and L1 distributions, we compared their densities. The Alus and exons were more abundant in the 3p21.3-p22 region whereas the L1s were more abundant in the 9q32 region. To test the significance of these differences, we applied a statistical analysis technique known as two-sample paired t-test. The significant difference (p<0.05) between the regions was observed in exon and Alu but not in L1 (Table 1).

Next, densities of exon, Alu and L1 in each 100k bp segment were plotted in Fig. 1. Due to extremely low gene content, no or little tendency was observed in the region 9q32. In 3p21.3-p22, positive correlation against exon density was observed for Alu(r = 0.42) and negative correlation against exon density was observed for L1 (r = -0.42). When the data in both regions were taken into

to 170k bp, the latter correlation coefficient also became a large negative value -0.7 (data not shown). account, the correlation coefficient between exon and Alu densities became 0.62 while that between exon and L1 densities was unchanged (r =-0.42). However, when the segment size was enlarged up

These results suggest that Aluelements but not LIs have a tendency to cluster into regions where the gene density is high. Although existence of direct relationships between exons and repetitive elements is not clear yet, the observed correlations might have an influence on gene function or gene expressions.

Table1:Comparise	exon, exon,	Atu and L1 Alu	Consity L1
3p21-22(1/k bp)	0.15	0.34	0.12
9q32(1/k bp)	0.02	0.16	0.16
Std.	0.12	0.13	0.11
t	3.28	4.78	-0.89
p	1.7×10^{-3}	4.5×10^{-5}	0.20



Figure 1: Correlation between exon density and Alu or L1 A, Correlation between Alu density and Exon density. Correlation coefficient is 0.62. B, Correlation between L1 density and exon density. Correlation coefficient is -0.42. Squares are 3p21.3-p22. Diamonds are in 9q32.

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