



Fig. 1 **a** Long-range polymerase chain reaction (LRP) of the SVA retrotransposon in X-linked dystonia-parkinsonism. *Lane 1*: Lambda DNA/*Hind*III marker; *Lanes 2 and 3*: normal individuals with wild allele; *Lanes 4, 6, and 7*: affected individuals with SVA retrotransposon in TAF1 gene; *Lane 5*: female carrier with both wild allele and SVA retrotransposon. The length of SVA retrotransposon and normal allele is 3,229 and 599 bp, respectively. **b** LRP-RFLP analysis using the

*Bam*HI. Digestion of LRP products obtained from an XDP patient shows two bands, 2,639 and 590 bp, as predicted. *C*, cut; *U*, uncut, *S*, Lambda DNA/*Hind*III marker. **c** Sequence analyses of the DSC3. Sequence fluorescent chromatograms show hemizyosity for a wild allele C in a healthy man (*upper panel*), hemizyosity for a changed allele T in an affected man (*middle panel*), and a heterozygous C/T in an obligate female carrier (*lower panel*)

39 (PCR) sequencing of several DSCs [2, 3]. Since 1991, 505
 40 XDP cases have been registered with the Philippine XDP
 41 project based at the Philippine Children's Medical Center in
 42 Metro Manila and in Roxas City, Capiz [1]. Improving
 43 diagnostic methods for the detection of nucleotide changes
 44 would contribute to enhance national surveillance of XDP in
 45 the Philippines and genetic counseling to XDP families.
 46 Therefore, conventional Southern analysis or direct PCR
 47 sequencing might be replaced by an alternative method
 48 using long-range polymerase chain reaction (LRP). This
 49 article describes the LRP protocol used to accurately detect
 50 the SVA retrotransposon.

51 Forty-two DNA samples were used from 14 unrelated
 52 normal controls, 20 XDP patients, and 8 obligate female
 53 carriers from 16 families residing in Panay, in that had been
 54 previously diagnosed with XDP by Southern analysis [2]. In
 55 addition, three DNA samples extracted from paraffin-
 56 embedded affected brain tissue were also used. LRP was
 57 performed as described in the supplementary conditions.
 58 CR-direct sequencing of DSC3 was also performed as de-
 59 scribed elsewhere [3]. As shown in Fig. 1, DNA fragments

with or without SVA retrotransposon were amplified (Fig. 1a). 60
 Restriction fragment length polymorphism analysis using the 61
 restriction enzyme *Bam*HI showed the predicted size of 62
 DNA fragments in the samples from XDP patients 63
 (Fig. 1b). The genotypes obtained by means of LRP were 64
 completely consistent with those previously determined by 65
 Southern analysis. Moreover, SVA retrotransposon is shown 66
 to be in complete linkage disequilibrium with allele T at the 67
 DSC3 (Fig. 1c). No false-positive results were obtained in 68
 controls. No amplification of the SVA retrotransposon was 69
 shown in DNA samples extracted from paraffin-embedded 70
 brain tissue, probably due to genomic DNA fragmentation. 71
 However, the direct sequencing of PCR products demon- 72
 strated the presence of the disease-associated allele T at 73
 the DSC3 in the pathologically confirmed XDP samples 74
 (data available upon request). These results prove that 75
 equivalent results can be obtained from the two methods. 76
 In conclusion, considering the multistep purification pro- 77
 cedures in PCR-direct sequencing and laborious techni- 78
 que in Southern analysis, LRP-based genetic testing 79
 might be preferred for the initial surveillance on XDP. 80

81 PCR-direct sequencing of the DSCs will further confirm
82 the results.
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Conflict of interest The authors declare that they have no conflict of
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101

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96 **Ethical standards** The study was performed according to a protocol
97 reviewed and approved by the Ethics Committee of the Graduate
98 School of Medicine, University of Tokushima.
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