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To cite this version:
Ariela Fundia, María Gorostiaga, Marta Mudry. Expression of common fragile sites in two Ceboidea species: Saimiri boliviensis and Alouatta caraya (Primates: Platyrhini). Genetics Selection Evolution, BioMed Central, 2000, 32 (1), pp.87-97. 10.1051/gse:2000108. hal-00894297

HAL Id: hal-00894297
https://hal.archives-ouvertes.fr/hal-00894297
Submitted on 1 Jan 2000

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Original article

Expression of common fragile sites in two Ceboidea species: *Saimiri boliviensis* and *Alouatta caraya* (Primates: Platyrhini)

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(Received 7 September 1998; accepted 22 November 1999)

Abstract – Fragile sites are points of preferential breakage that may be involved in chromosome rearrangements. Induction of common fragile sites (c-fra) and spontaneous breakage were analyzed in two New World Monkeys species: *Saimiri boliviensis* (SBO) and *Alouatta caraya* (ACA). Spontaneous chromosome aberrations were analyzed on untreated lymphocyte cultures with Brögger’s formula (1977). SBO presented a low level of spontaneous breakage, while higher frequencies were detected in ACA in which bands 1q23; 2q13 and 11q19 were significantly affected \( (p < 0.01) \). The populational distribution of c-fra was analyzed by the Chi\(^2\) test in FUDR plus caffeine treated cultures. A total of 21 c-fra was identified in SBO and 24 in ACA. Fragile sites A\(_1\)q33, B\(_1\)p21, B\(_4\)p14, C\(_3\)q23 and C\(_5\)q22 were identified in all analyzed SBO specimens. The most frequent c-fra identified in ACA specimens were 1q22, 1q31, 2q22, 8q14, 12q31, 13q22, 14q15 and Xq22. Fragile sites A\(_1\)q31, A\(_1\)q33, B\(_1\)q14, B\(_3\)q13, B\(_4\)q21 and Xq22 identified in SBO and 1q31, 1q33, 2q22, 4q21, 6q13, 13q22 and Xq22 from ACA were the most conserved sites. A low coincidence between the location of c-fra and that of heterochromatin and breakpoints involved in euchromatic rearrangements known for these genera, was established.

Ceboidea / fragile sites / chromosomal rearrangements / heterochromatin / evolution

Résumé – Sites communs de fragilité chromosomique chez deux espèces de Ceboïdes : *Saimiri boliviensis* et *Alouatta caraya* (Primates : Platyrhini). Les sites fragiles sont des régions de cassure préférentielle, dans le génome, qui peuvent être associées à des remaniements chromosomiques. On a étudié les sites communs fragiles (c-fra) et les cassures spontanées dans deux genres de primates du Nouveau Monde : *Saimiri boliviensis* (SBO) et *Alouatta caraya* (ACA). Les cassures spontanées

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ont été étudiées avec la formule de Brögger (1977) sur des cultures de lymphocytes non traités. Alors que SBO présente très peu de cassures spontanées, celles-ci sont fréquentes chez ACA, où trois bandes : 1q23, 2q13 et 11q19 sont significativement affectées \((p < 0,01)\). La distribution spécifique des c-fra a été analysée par le test du Chi² dans des cultures traitées par le FUdR et la cafétine. Un total de 21 c-fra chez SBO et 24 chez ACA a été observé. Les c-fra A1q33, B1q21, B3q14, C5q23 et C5q22 ont été identifiées dans tous les spécimens SBO. Les c-fra les plus fréquents identifiées chez ACA sont 1q23, 1q31, 1q33, 2q22, 8q14, 12q31, 13q22, 14q15 et Xq22. Les c-fra les plus conservés chez SBO sont A1q31, A1q33, B1q14, B3q13, B4q21 et Xq22 et chez ACA 1q31, 1q33, 2q22, 4q21, 6q13, 13q22 et Xq22. Nous avons établi une faible coïncidence entre l’emplacement des c-fra et celui de l’hétérochromatine et des cassures impliquées dans les remaniements de l’euchromatine connus dans ces genres.

Ceboides / sites fragiles / remaniements chromosomiques / hétérochromatine / évolution

1. INTRODUCTION

Chromosomal fragile sites are points on chromosomes which show non-random gaps or breaks under specific conditions. They are classified as rare (carried by few individuals) or common (virtually in all individuals) and are subdivided according to the conditions used for their induction. Rare folate sensitive fragile sites are expansions (dynamic mutations) of CCG-repeat sequences. Rare FRA16B and FRA10B sites are expansions of very AT-rich minisatellites. In contrast, sequence data for common sites show no striking features such as trinucleotide or minisatellite repeats [63]. These sites have been shown to display a number of characteristics of unstable and highly recombinogenic DNA in vitro, including chromosome rearrangements, sister chromatid exchanges and intrachromosomal gene amplifications [26]. Although there has been substantial advancement in the study of their molecular structure, the potential for a relationship with disease is still unknown except for the association with mental retardation (fragile X syndrome) [63]. Autosomal fragile sites have been related to the origin of constitutional or cancer rearrangements [23, 33, 49, 61, 63] and they can be targets of mutagens and carcinogens [2, 3, 71]. They have also been associated with chromosomal changes during evolution [6, 28, 31, 38, 39].

Examples of chromosomal variation are common in primate speciation [6, 13, 65], pericentric inversions and heterochromatic block variations being more frequently observed [48]. Chromosome comparison using banding methods and in situ hybridization has demonstrated that human chromosomes show a high homology with some Platyrrhini species [7–9, 43, 53] as well as other anthropoid species [6, 48, 57] and other mammals [68]. Taking into account that chromosomal rearrangements could be incorporated in the course of the evolutionary process, comparative cytogenetics has been used in phylogenetic studies [6, 8, 13, 51]. Considering that common fragile sites (c-fra) are reliable markers of genetic instability [24], fragile site studies provide a widely applicable means to evaluate the change in chromosome structure and its possible implications in speciation. In the present work, we provide new evidence on primate chromosome variability, evaluating spontaneous breakage and distribution of common fragile sites in two Ceboida species.
2. MATERIALS AND METHODS

C-fra expression was analyzed in heparinized peripheral blood samples from 16 specimens: 12 ACA and 4 SBO. Two cultures were set up simultaneously for each specimen in F10 medium with phytohemagglutinin M (0.1 μg·mL⁻¹, SIGMA) and fetal bovine serum (5%, GIBCO) for 72 h at 37 °C. Spontaneous breakage was analyzed in an untreated culture (control) and fragile sites were induced by known fragile site inducers such as fluorodeoxyuridine (FUdR) (10 μg·mL⁻¹) for the final 24 h of culture [20] and caffeine (2.2 mM) for the last 6 h [70].

Cells were routinely harvested and 25 to 70 Giemsa stained metaphases were analyzed on coded slides to record the presence of chromosome aberrations (CA), following the “International System for Human Cytogenetic Nomenclature” [29]. Slides with abnormal cells were destained and re-analyzed after sequential G-banding [59] to identify the breakpoints involved in CA. The karyotype of each species was considered following previously published works [21, 22, 25, 45–47]. Spontaneous CA were analyzed with Brögger’s formula (1977) [4], and fragile sites were defined by the Chi² test with Yates correction [27]. A haploid karyotype was considered for both statistical analyses assuming that all bands had an equal probability of breakage.

3. RESULTS

The most frequent CAs observed were gaps and breaks, while a low proportion of acentric fragments or triradial figures was only found in treated cultures. Chromosome or chromatid gaps, breaks and acentric fragments were scored as single chromosome events and dicentric or triradial configurations as two chromosome events. G-band analysis of control cultures allowed the identification of 6 and 39 spontaneous CAs in SBO and ACA specimens, respectively. Based on a SBO haploid karyotype of 257 bands [25, 46], statistical analysis with Brögger’s formula showed that any band with 2 or more lesions was non-randomly damaged (p < 0.0005). Since these six aberrations were located on different bands, no bands significantly involved in spontaneous breakage were found in SBO. On the other hand, spontaneous breakage analysis in ACA considering a haploid karyotype of 287 bands [21, 22, 46, 47] demonstrated that any band with 2 or more lesions was non-randomly damaged (p < 0.01), identifying 3 bands: 1q23, 2q13 and 11q19 which were hot-spots for spontaneous breakage.

A total of 245 and 328 CAs was identified with sequential G-banding in SBO and ACA treated cultures, respectively. Based on a SBO haploid karyotype, the expected number of breaks per band for the 245 observed aberrations is 0.95. Chi² analysis showed that any band with five or more lesions is non-randomly damaged in excess (p < 0.001), indicating 21 induced fragile sites (Tab. I). The expression frequencies of these fragile sites confirmed that all sites were common (c-fra). Five of the 21 fragile sites (24%), located at A1q33, B1p21, B4p14, C3q23 and C5q22 were identified in all SBO specimens, 13 sites (62%) were induced in 3 specimens and only 3 sites (14%) were detected in 2 specimens. SBO specimens exhibited 13 to 19 of the 21 c-fra (Tab. I). Some of these fragile sites are shown in Figure 1. Based on an ACA haploid karyotype,
Table I. Numbers of chromosome aberrations and common fragile sites induced in *Saimiri boliviensis*.

| Fragile sites | Chromosome aberrations in each specimen |
|---------------|----------------------------------------|
|               | 1 | 2 | 3 | 4 | Total |
| A<sub>1</sub>q21 | 2 | 0 | 3 | 4 | 9   |
| A<sub>1</sub>q31 | 3 | 1 | 1 | 0 | 5   |
| A<sub>1</sub>q33 | 1 | 1 | 1 | 2 | 5   |
| A<sub>1</sub>q35 | 2 | 0 | 2 | 1 | 5   |
| A<sub>2</sub>q12 | 3 | 5 | 1 | 0 | 9   |
| B<sub>1</sub>p21 | 4 | 3 | 1 | 1 | 9   |
| B<sub>1</sub>q14 | 5 | 4 | 0 | 1 | 10  |
| B<sub>1</sub>q23 | 1 | 0 | 3 | 1 | 5   |
| B<sub>2</sub>q21 | 0 | 0 | 2 | 3 | 5   |
| B<sub>2</sub>q23 | 0 | 2 | 3 | 0 | 5   |
| B<sub>2</sub>q31 | 3 | 2 | 3 | 0 | 8   |
| B<sub>2</sub>q33 | 5 | 6 | 0 | 1 | 12  |
| B<sub>3</sub>q13 | 4 | 2 | 1 | 0 | 7   |
| B<sub>3</sub>q15 | 3 | 2 | 1 | 0 | 6   |
| B<sub>4</sub>p14 | 1 | 2 | 1 | 1 | 5   |
| B<sub>4</sub>q21 | 4 | 0 | 4 | 1 | 9   |
| B<sub>4</sub>q23 | 3 | 2 | 3 | 0 | 9   |
| C<sub>3</sub>q23 | 5 | 1 | 1 | 1 | 8   |
| C<sub>3</sub>q25 | 2 | 2 | 0 | 1 | 5   |
| Xq22          | 2 | 2 | 1 | 6 | 5   |
| Total number of c-fra | 57 | 38 | 32 | 19 | 146 |
| Number of CA recorded | 84 | 74 | 56 | 31 | 245 |
| Number of cells analyzed | 70 | 50 | 50 | 50 | 220 |
| Number of abnormal cells | 50 | 36 | 29 | 20 | 135 |

Figure 1. G-banded chromosomes showing fragile sites (--) induced in SBO specimens.
the expected number of breaks per band for the 328 aberrations observed is 1.14. Statistical analysis showed that any band with five or more lesions is significantly damaged \((p < 0.005)\), identifying 24 c-fra (Tab. II). Fourteen out of 24 c-fra were fragile in 50–83% of the ACA population. The most frequent fragile site was 2q22 (Fig. 2) expressed in 10 specimens (83%), while 4q21, 9p13, 9q13, 11q13 and 15q21 were observed in only six specimens (50%). ACA specimens exhibited 5 to 19 of the 24 fragile sites.

Locations of fragile sites were compared with heterochromatic regions and breakpoints involved in euchromatic rearrangements known at present. No chromosome rearrangements were found in the SBO specimens presently analyzed nor in our previous studies with different specimens. A heterochromatic polymorphism at chromosome B11p14 was observed in these specimens, but no fragile sites were found at this site. Only 3/21 (14%) of the c-fra sites (B1q14, B1q23, B4p14) coincided with C-bands in SBO. No relationship between

| Common Fragile sites | Chromosome aberrations in each specimen |
|----------------------|----------------------------------------|
|                      | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | Total |
| 1q13                 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 3 | 0 | 1 | 6 |
| 1q23                 | 1 | 0 | 0 | 2 | 2 | 1 | 0 | 1 | 1 | 1 | 1 | 5 | 15 |
| 1q31                 | 2 | 1 | 0 | 2 | 0 | 1 | 0 | 2 | 0 | 2 | 0 | 2 | 12 |
| 1q33                 | 1 | 0 | 0 | 4 | 0 | 0 | 0 | 1 | 1 | 1 | 1 | 1 | 10 |
| 2q13                 | 0 | 0 | 0 | 1 | 0 | 2 | 0 | 1 | 0 | 1 | 5 | 0 | 10 |
| 2q22                 | 1 | 1 | 1 | 1 | 3 | 2 | 0 | 1 | 1 | 2 | 0 | 1 | 14 |
| 2q36                 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 2 | 5 |
| 3q31                 | 0 | 0 | 0 | 2 | 0 | 2 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 6 |
| 4q21                 | 0 | 0 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 1 | 0 | 1 | 6 |
| 6q15                 | 0 | 1 | 0 | 0 | 0 | 2 | 0 | 0 | 1 | 0 | 2 | 1 | 7 |
| 8q14                 | 1 | 3 | 0 | 2 | 0 | 2 | 2 | 1 | 0 | 0 | 3 | 5 | 19 |
| 9q13                 | 1 | 2 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 3 | 2 | 10 |
| 9q13                 | 3 | 1 | 0 | 2 | 0 | 2 | 0 | 0 | 1 | 1 | 0 | 0 | 10 |
| 11q13                | 1 | 0 | 0 | 0 | 1 | 3 | 0 | 1 | 0 | 0 | 1 | 1 | 8 |
| 11q19                | 1 | 0 | 0 | 0 | 2 | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 6 |
| 11q23                | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 1 | 0 | 0 | 3 |
| 12q13                | 1 | 1 | 0 | 3 | 0 | 0 | 1 | 0 | 2 | 1 | 0 | 1 | 10 |
| 13q13                | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 0 | 2 | 1 | 0 | 0 | 6 |
| 13q22                | 1 | 0 | 0 | 2 | 1 | 0 | 0 | 1 | 1 | 2 | 0 | 2 | 10 |
| 13q24                | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 2 | 1 | 5 |
| 14q15                | 0 | 0 | 1 | 1 | 0 | 1 | 1 | 1 | 0 | 1 | 0 | 1 | 7 |
| 15q21                | 0 | 1 | 0 | 1 | 1 | 1 | 0 | 0 | 0 | 2 | 1 | 0 | 7 |
| 16q13                | 0 | 2 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 6 |
| Xq22                 | 1 | 0 | 1 | 2 | 2 | 1 | 1 | 0 | 1 | 1 | 0 | 1 | 11 |

Total number of c-fra: 15 13 5 29 14 20 9 17 14 22 20 34 212
Number of CA recorded: 26 24 14 41 20 26 17 28 20 25 34 53 328
Number of cells analyzed: 25 25 50 50 50 25 25 25 25 25 50 400
Number of abnormal cells: 14 16 12 14 16 27 17 18 18 17 16 38 223
induced c-fra sites and chromosomal changes known in SBO was found. In ACA, a particular sex determination system was observed, resulting from a reciprocal translocation t(7;Y), but no heterochromatic region or fragile site were observed on these chromosomes. Only c-fra 1q31 from ACA coincided with a breakpoint involved in a pericentric inversion proposed by Mudry et al. [43], demonstrating a low coincidence (1/24, 4%) with rearranged sites. Considering the total number of fragile sites identified in each species, no significant correlation was found between heterochromatic regions or structural changes and fragile sites.

4. DISCUSSION

It has been suggested that fragile sites are regions susceptible to breakage and rearrangements that could be involved in chromosome evolution. Many reports have established a theoretical correlation between the location of human fragile sites and bands involved in rearrangements during primate chromosomal evolution [5, 6, 39, 60]. The possible mechanisms and pathway of karyotype evolution in primates have been extensively discussed [12, 13, 47, 52]. The immense variety of karyotypes of extant forms provides suggestive evidence that chromosome change has played and continues to play a major role in
Fragile sites expression in Ceboidea  

evolution [11, 12, 14–16, 30]. What is the relationship between chromosome change and evolutionary change? The ability to determine accurately the type and number of rearrangements is a critical step in understanding chromosomal evolution [1].

Spontaneous breakage has rarely been described in the karyotypes of pri- 

mates up to now and to our knowledge there are no other reports of spontan- 

eous chromosome fragility in different Ceboidea species. A low frequency of spontaneous CA was found in SBO specimens (6%), while ACA presented higher levels (30%). Different frequencies of spontaneous breakage have been previously reported in other mammals, ranging from as few as 8% to as many 64% [2, 3, 38, 60, 69]. In addition, three spontaneous fragile sites, coincident with induced ones, were found to be significantly damaged in ACA, suggesting that these areas are more susceptible to chromosomal breakage. The specific in- 

volvement of certain bands in spontaneous breakage from New World Monkeys provides new evidence for the particular variability of the Ceboidea genome.

Relatively few reports have been published on induction of fragile sites in species other than man [17, 18, 34, 38, 54–56, 62, 64]. C-fra induction was reported in a few neotropical primates: *Cebus apella* [19, 41] and *Alouatta caraya* [21]. Induction of fragile sites was also described in gorilla, chimpanzee and orangutan, showing an evolutionary conservation of these sites between the Great Apes and man and suggesting that fragile sites have been highly conserved during primate evolution [58, 60, 70]. In order to analyze the con- 

servation of fragile sites in Ceboidea, the present data were compared to 

previous results on *Cebus apella paraguayanus* (CAP) chromosomes identi- 

fying 11 induced fragile sites (2q13, 2q26, 3q31, 5q22, 6q21, 11q15, 12q22, 19q13, 19q22, 20q13 and Xq22) [22, 46]. Taking into account the chromosome homologies previously described in SBO, ACA and CAP [43], a homologous c-fra at band Xq22 was observed in all three species. This finding is in agreement with the well-known conservation of the X chromosome [12, 43, 57, 67, 68]. The three species also conserved a c-fra at bands 3q31 from CAP, B1q14 from SBO and 2q22 from ACA. C-fra A1q31, A1q33, B3q13 and B4q21 from SBO were homologous to c-fra 1q31, 1q33, 6q13 and 4q21 from ACA, respectively. C-fra 20q13 from CAP was homologous to 13q22 from ACA. These data are in agreement with our previous data on the variability of Ceboidea karyotypes [25, 43–47, 50, 52] and on those reported by other authors [9–11, 36].

It has been proposed that interbands between euchromatic and heterochro- 

matic regions are probably more susceptible to breakage [66]. A poor relation- 

ship was established between fragile site location and heterochromatic regions or breakpoints involved in euchromatic rearrangements known for CAP, SBO and ACA [5, 25, 32, 35–37, 40, 42, 43, 47, 52]. Three CAP c-fra sites (6q21, 11q15 and 12q22) coincide with heterochromatic bands. One of them, 12q22, is associated with a paracentric inversion involving the heterochromatic region observed in different *Cebus apella ssp* (Mudry, unpublished data), also reported for other heterochromatic regions [50]. Another c-fra, 11q15, coincides with a terminal deletion of a C-band in *C. a. nigritus* [44] and paracentric inversions observed in *C. a. robustus* and *C. a. xanthosternos* [37]. Three SBO c-fra sites (B1q14, B1q23, B4p14) are located at C-bands. Only one ACA c-fra (1q31) is located at a breakpoint of a pericentric inversion involved in the evolution of the Ceboidea karyotype [43]. Smeets and Klundert induced fragile site expression on
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chromosomes of the Great Apes and showed that 35% of fragile sites coincided with rearranged sites in primates [60]. In addition, a theoretical relationship was found between breakpoints in Ceboidea and human fragile sites, suggesting that the location of latent centromeres in Platyrrhini and heterochromatic regions were related [5]. Our results demonstrate that no important correlation exists between heterochromatin or structural changes and fragile sites induced in these Ceboidea species. In fact, to clarify the role of fragile sites in karyological evolution, it will be important to identify more fragile sites in a great number of individuals from different species, characterize chromosomal homologies between these species, identify more chromosome rearrangements involved in evolutionary pathways, map the breakpoints and compare them to fragile site locations.

ACKNOWLEDGEMENTS

We thank J. Ruiz for supplying SBO blood samples from CAPRIM (Argentine Primate Center); Dr. G. Zunino for his valuable work in the Argentinean forest supplying ACA and CAP samples; Lic. A. Delprat for her help with karyotype preparations; Mr. E. Crocito and R. Fraiman for their photographic work. This publication was produced during the UBACYT EX 288 Project.

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