Sporadic Premature Aging in a Japanese Monkey: A Primate Model for Progeria

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Abstract

In our institute, we have recently found a child Japanese monkey who is characterized by deep wrinkles of the skin and cataract of bilateral eyes. Numbers of analyses were performed to identify symptoms representing different aspects of aging. In this monkey, the cell cycle of fibroblasts at early passage was significantly extended as compared to a normal control. Moreover, both the appearance of senescent cells and the deficiency in DNA repair were observed. Also, pathological examination showed that this monkey has poikiloderma with superficial telangiectasia, and biochemical assay confirmed that levels of HbA1c and urinary hyaluronan were higher than those of other (child, adult, and aged) monkey groups. Of particular interest was that our MRI analysis revealed expansion of the cerebral sulci and lateral ventricles probably due to shrinkage of the cerebral cortex and the hippocampus. In addition, the conduction velocity of a peripheral sensory but not motor nerve was lower than in adult and child monkeys, and as low as in aged monkeys. However, we could not detect any individual-unique mutations of known genes responsible for major progeroid syndromes. The present results indicate that the monkey suffers from a kind of progeria that is not necessarily typical to human progeroid syndromes.

Introduction

Symptoms representing various aspects of aging at an early age are characteristic of progeria in humans. Progeria is a series of very rare genetic syndromes that are related to mutations in specific genes. For example, genes responsible for Werner syndrome (WS), Rothmund-Thomson syndrome (RTS), Bloom syndrome (BS), and Hutchinson-Gilford progeria syndrome (HGPS) are RECQ3 (WRN), RECQL4, RECQ2, and LMNA, respectively [1,2]. As summarized in Table 1, these syndromes are accompanied by common or separate physical abnormalities that can be ascribed to senile changes. An animal model is a powerful tool to investigate possible mechanisms underlying not only premature aging, but also normal aging. However, no useful animal models have so far been available. Although mice lacking the RECQL1 gene have been reported, their pathological changes are quite limited, probably due to a species difference between rodents and humans [3]. Thus, a primate model for progeria, which is taxonomically akin to humans and has a longer life-span than a rodent model, is needed to promote aging research. Recently, we have found a child Japanese monkey who displayed deep skin wrinkles and bilateral cataract at the age of one year. In the present study, we examined whether this monkey is actually progeroid or not from various viewpoints.

Methods

Animals and ethics statements

All animal experiments were planned and executed in strict concordance with Guide for the Care and Use of Laboratory Animals (8th Ed, 2011, The National Academies, USA) and Guidelines for Care and Use of Nonhuman Primates (Ver. 3, 2010, Primate Research Institute, Kyoto University). The protocol was approved by the Animal Welfare and Animal Care Committee, Primate Research Institute, Kyoto University (Permission No. 2012-115). All monkeys used in this study, weighing less than 10 kg, were kept in indoor individual cages (0.89 m×0.63 m×0.82 m) of the Primate Research Institute on a 12-h on/12-h off lighting schedule. In each cage, the monkeys were given ad libitum access to food and water, and a chain-hung wood block as a toy. No monkeys were sacrificed. Monkey N416, a female Japanese monkey (Macaca fuscata) who was supposed to be progeroid, was born in an outdoor cage (5 m×5 m×2.7 m, 6–7 monkeys per cage) and moved to an indoor individual cage at the age of one year and four months to avoid progress of cataract. Unfortunately, N416 suddenly died of bloat at the age of three years.
Measurements of cell proliferation

Cell proliferation was analyzed using fibroblast cells prepared from cultures of a tiny block of the ear skin as described previously [4]. To determine the duplication time of each cell, cultures were made using a medium of Amnio Max II Complete (Gibco BRL, USA) in a 60×15 mm petri dish for cell culture (FALCON, 353002, USA). After appropriate proliferation of cells in the primary dish culture, cells were harvested, and then cell suspension (25–30 cells/mL) was transferred to a 35×6×10 mm cell culture dish (Advanced glass bottom, Greiner Bio-one, 627965, Germany). One day later, recordings of the cell division were performed every five minutes with a time-lapse imaging system (Cell Observer, Carl Zeiss, Germany). The time of a cell cycle was figured out by chasing each cell division based on imaging data collected in 70-hr cultures: the time was calculated as an interval time from the first division to the second division of each cell. A total of 44 or 47 cells were analyzed in monkey N416 and a normal control, respectively. To measure the population growth activity, fibroblast cells derived from monkey N416 and normal infant and aged monkeys (all at passage 8) were seeded at 1×10^5 cells/well in gelatinized 6-well plates with DMED containing 15% FBS, 0.1 mM nonessential amino acids, 2 mM L-glutamine, 1 mM sodium pyruvate, 0.11 mM 2-mercaptoethanol, 100 U/ml penicillin, and 100 μg/ml streptomycin. The total cell numbers in three independent wells were counted every 3 days up to day 12.

Detection of senescent cells

Senescent cells were detected with ß-galactosidase assay using fibroblast cells prepared from monkey N416, a normal infant and a normal aged monkey. The number of passage of fibroblast cells derived from monkey N416 and the infant and the aged controls was 5, 7, and 5, respectively. Senescence-associated ß-galactosidase activity was detected by using Cellular Senescence Assay Kit (CBA-230, Cell Biolabs, USA). Cells were fixed and incubated with X-gal overnight. Five images were randomly captured (BZ-9000, Keyence, Japan) from each culture dish of the three monkeys, and the numbers of stained and unstained cells were counted.

Quantification of DNA repair

To test DNA repair deficiency in monkey N416, we quantified DNA damage by counting apurinic/apyrimidinic sites (AP sites), which are the target of base excision repair. DNA was freshly prepared with DNeasy Blood & Tissue Kit (69504, Qiagen, USA) from fibroblast cells of the three monkeys used for the senescent cell detection. Fibroblast cells from the infant control had been

Table 1. Physical signs and symptoms characteristic of human progeroid syndromes and monkey N416.

| Signs and symptoms                              | WS | RTS | HGPS | BS | N416 |
|------------------------------------------------|----|-----|------|----|------|
| cataract                                       | ✓  | ✓   | ✓    | ✓  | ✓    |
| abnormal glucose metabolism                    | ✓  |     | ✓    | ✓  | ✓    |
| urinary hyaluronan                              | ✓  | ✓   | ✓    | ✓  | ✓    |
| decline in proliferative potency of fibroblasts|     | ✓   | ✓    | ✓  | ✓    |
| at early passage                               | ✓  | ✓   | ✓    | ✓  | ✓    |
| increase in senescent fibroblast cells         | ✓  | ✓   | ✓    | ✓  | ✓    |
| DNA repair deficiency                          | ✓  | ✓   | ✓    | ✓  | ✓    |
| poikilodermia                                  | ✓  |     | ✓    | ✓  | ✓    |
| short stature, low bodyweight                  | ✓  | ✓   | ✓    | ✓  | ✓    |
| facial proportion                              | ✓  | ✓   | ✓    | ✓  | ✓    |
| loss of hair                                   | ✓  | ✓   | ✓    | ✓  | ✓    |
| facial rash                                    | ✓  | ✓   | ✓    | ✓  | ✓    |
| intractable skin ulcers                        | ✓  |     | ✓    | ✓  | ✓    |
| prominent scalp vein                           | ✓  |     | ✓    | ✓  | ✓    |
| sun-sensitivity                                | ✓  | ✓   | ✓    | ✓  | ✓    |
| skeletal defects                                | ✓  | ✓   | ✓    | ✓  | ✓    |
| abnormal dentation                             | ✓  |     | ✓    | ✓  | ✓    |
| characteristic voice                           | ✓  | ✓   | ✓    | ✓  | ✓    |
| soft-tissue calcification                      | ✓  | ✓   | ✓    | ✓  | ✓    |
| contracture of joints                          | ✓  |     | ✓    | ✓  | ✓    |
| atherosclerosis                                | ✓  | ✓   | ✓    | ✓  | ✓    |
| malignant tumors                               | ✓  | ✓   | ✓    | ✓  | ✓    |
| hypogonadism                                   | ✓  | ✓   | ✓    | ✓  | ✓    |
| immunodeficiency                               | ✓  | ✓   | ✓    | ✓  | ✓    |
| onset after adolescence                        | ✓  | ✓   | ✓    | ✓  | ✓    |
| onset within one year of age                   | ✓  | ✓   | ✓    | ✓  | ✓    |

BS: Bloom syndrome, HGPS: Hutchinson-Gilford progeria syndrome, nt: not tested, RTS: Rothmund-Thomson syndrome, WS: Werner syndrome, (√): Controversial [10,11].

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anesthetized with ketamine hydrochloride (5 mg/kg b.wt., i.m.) with a 0.3 T apparatus (Vento, Hitachi). The monkeys were age (2 years old), and an aged monkey (28 years old) were taken

supramaximal amplitude.

abductor digiti minimi and the ulnar nerve was stimulated 2–3 cm joints of the little finger. To measure the motor conduction electrodes attached to the distal and proximal interphalangeal supramaximal stimuli were delivered through a pair of ring proximal to the elbow and 2–3 cm proximal to the wrist, and surface electrodes were placed over the ulnar nerve 2–3 cm the measurement. The room temperature was kept at 20

Statistics

Measurements of blood and urinary biomarkers

Blood and urine samples were obtained from monkey N416 and a control monkey of the same age (2 years old) were obtained with Asteion TSX-021B (Toshiba Medical Systems). The monkeys were anesthetized with ketamine hydrochloride (5 mg/kg, i.m.) and medetomidine (100 μg/kg, i.m.), followed by head fixation with a stereotaxic apparatus [5]. Three-dimensional reconstruction was done with VirtualPlace (Aze).

X-ray CT and MRI

Images of the head and extremities of monkey N416 and a control monkey of the same age (2 years old) were taken with a 0.3 T apparatus (Vento, Hitachi). The monkeys were anesthetized with ketamine hydrochloride (5 mg/kg b.wt., i.m.) and medetomidine (100 μg/kg b.wt., i.m.), followed by head positioning with the stereotaxic apparatus [5].

Nerve conduction velocity

To assess the functional viability of the peripheral nervous system, the conduction velocity of sensory and motor nerves was measured from the ulnar nerve of monkey N416, in comparison with those in control Japanese monkeys including 5 child monkeys (Child; 2 years old), 5 adult monkeys (Adult; 5–8 years old), and 5 aged monkeys (Aged; 25–29 years old). Measurements were done with NeuroPak (Nihon Kohden, Tokyo). The monkeys were sedated with ketamine hydrochloride (2.5 mg/kg) and medetomidine (100 μg/kg), and anesthetized with 2% sevoflurane during the measurement. The room temperature was kept at 20°C. To measure the sensory conduction velocity (SCV), pairs of recording surface electrodes were placed over the ulnar nerve 2–3 cm proximal to the elbow and 2–3 cm proximal to the wrist, and supramaximal stimuli were delivered through a pair of ring electrodes attached to the distal and proximal interphalangeal joints of the little finger. To measure the motor conduction velocity (MCV), recording surface electrodes were placed over the abductor digitii minimi and the ulnar nerve was stimulated 2–3 cm proximal to the elbow and 2–3 cm proximal to the wrist, with supramaximal amplitude.

Measurements of blood and urinary biomarkers

Blood and urine samples were obtained from monkey N416 (2 years and 6 months old), 5 child monkeys (age-matched), 5 adult monkeys (6–9 years old), and 5 aged monkeys (21–27 years old). Measurements of hemoglobin A1c (HbA1c; blood), hyaluronan (urine and serum), glucose (urine and serum), low-density lipoproteins (LDL; serum), high-density lipoproteins (HDL; serum), and triglyceride (TG; serum) were carried out by a commercial test company (Falco Biosystems).

Statistics

Student’s t-test was applied for cell proliferation time comparison. One-way ANOVA and post hoc test (Bonferroni) were applied for analyses of the ratio of senescent cells, the number of AP sites, the levels of biomarkers, and the conduction velocity of peripheral nerves (Kaleidagraph 4.1, Hulinks, Japan). Linear discriminant analysis was also applied for the peripheral nerve conduction velocity.

DNA sequence

The sequences of candidate genes for progeroid syndromes, RECQ3 (Werner syndrome), RECQL4 (Rothmund-Thomson syndrome), RECQ2 (Bloom syndrome), and LMNA (Hutchinson-Gilford progeria syndrome), were determined by standard PCR and the direct sequence protocol with ExTaq DNA polymerase (Takara Bio Inc., Shiga, Japan) and BigDye Terminator v3.1 Cycle Sequencing Kit with a 3130 Genetic Analyzer (Applied Biosystems, CA, USA). Primers used in PCR are shown in Table S1. Furthermore, to identify possible genetic causes specifically occurring in monkey N416, broader candidate gene approaches were performed in 100 macaque monkeys including monkey N416 with Next Generation Sequencer (NGS) of HiSeq2000 (Illumina Inc., CA, USA). The candidates for progeroid-related genes were listed in Table S2. Genomic DNA samples were obtained from 100 macaque monkey blood samples including monkey N416. Each library for the NGS analysis was constructed by using the standard Illumina TruSeq DNA Library Prep protocol.

Results

Physical appearance

Two major appearances characteristic of monkey N416 were deep wrinkles of the skin and cataract on bilateral sides. The skin wrinkles have been eminent since she was a baby, while such wrinkles were not observed in her mother (Fig. 1A). As monkey N416 grew up, her wrinkles seemed to gradually decrease but still remained at the age of two years (Fig. 1B). Indeed, X-ray CT clearly revealed the wrinkled skin of this monkey (Fig. 1C; imaged later than the time when Fig. 1B was taken) and, as a control, the smooth skin of an age-matched normal monkey (Fig. 1D). There was no sign of ulceration of the skin, and hyperkeratosis was slightly observed in the extremities. Cataract occurred in one eye as early as monkey N416 was 9 months old, and progressed bilaterally as she grew around two years old (Fig. 1B). Based on the fact that this monkey usually reached for primate pellets in a box attached to her own cage and smelled them before eating, it appeared that her vision was still left but was only weak. When we examined her under anesthesia, we could not find out either rigidity or hyperflexibility of the limbs. No skeletal anomaly, osteoporosis, or dental crowding was found in monkey N416.

Fibroblast cell cycle

Our time-lapse video analysis at passage 5 confirmed that the time for fibroblast cell cycle was significantly longer in the model monkey N416 than in a normal control (Fig. 2A; N416 28.8±6.8 hr, control 20.0±4.6 hr, t = −7.17, p < 0.01). Population growth of fibroblast cells at passage 8 from the aged monkey reached a plateau at day 6. On the other hand, population growth of fibroblast cells from monkey N416 and the infant monkey continued up to day 12 (Fig. 2B).

Senescent cells

Cytotoxic assay of β-galactosidase activity revealed that the ratio of senescent (β-galactosidase-positive) fibroblast cells was higher in the aged monkey than in the infant monkey. Their ratio in monkey N416 was significantly lower than in the aged monkey and higher than in the infant monkey (Fig. 2G; aged 19.6±2.8%, 28.8±6.8 hr, control 20.0±4.6 hr, t = −7.17, p < 0.01). Population growth of fibroblast cells at passage 8 from the aged monkey reached a plateau at day 6. On the other hand, population growth of fibroblast cells from monkey N416 and the infant monkey continued up to day 12 (Fig. 2B).

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Premature Aging in Nonhuman Primates

DNA repair
Deficiency of DNA repair was estimated through AP site quantification of DNA extracted from fibroblast cells. One-way ANOVA and post hoc test revealed that numbers of AP sites were higher in cells from monkey N416 and UV-irradiated cells from the infant monkey than in cells from the aged and infant monkeys (Fig. 2D; aged 12.2±1.2, infant 14.6±1.7, UV-irradiated infant 22.3±1.4, N416 19.4±1.2, F[3,8]=33.2, p<0.001, one-way ANOVA; N416 vs. aged, UV-irradiated infant vs. aged, UV-irradiated infant vs. infant, p<0.01, N416 vs. infant, p<0.05, post hoc test).

Blood and urinary biomarkers
We measured levels of blood and urinary biomarkers in monkey N416 and compared these values with those in aged, adult, and child monkeys. HbA1c in monkey N416 was not different from those in the aged and child groups (one-way ANOVA and post hoc test). Likewise, the motor conduction velocity (MCV) in monkey N416 was measured from the ulnar nerve. The sensory conduction velocity (SCV) in monkey N416 was compared with that in aged, adult, and child monkeys; each value was 62.1, 61.2±4.5 (n=5), 63.2±4.3 (n=5), and 57.0±13.8 (n=5) ms, respectively (Fig. 5A). It was revealed that the SCV in monkey N416 and the aged group was significantly slower than in the adult and child groups (one-way ANOVA and post hoc test). Likewise, the motor conduction velocity (MCV) in monkey N416 was examined in comparison with that in the aged, adult, and child groups; each value was 62.1, 61.2±4.5 (n=5), 63.2±4.3 (n=5), and 57.0±13.8 (n=5) ms, respectively (Fig. 5B). Thus, the MCV in monkey N416 was not significantly different from that in the aged, adult, or child monkey group (one-way ANOVA). Scattergram of SCV and MCV shows that data points for the aged, adult, and child monkeys appeared to constitute three separate clusters, and the data point for monkey N416 was included within the cluster for the aged group (Fig. 5C).

DNA sequences of progeroid-related genes
The sequences of progeroid-related genes, RECQ3, RECQ14, RECO2, and LMNA, were determined in monkey N416 and normal Japanese monkeys (n = 7, 41, 2, and 4, respectively) and a normal rhesus monkey, as referenced with those of rhesus monkey genomic sequence (rheMac2). Though there were five non-synonymous point mutations in recQ14 gene of monkey N416 with reference to rhesus sequences (exon3 183A>G, 387C>T, exon4 157G>A, 208T>C, 315C>G), such mutations were not significantly different from that in the aged, adult, or child monkey group (one-way ANOVA). Scattergram of SCV and MCV shows that data points for the aged, adult, and child monkeys appeared to constitute three separate clusters, and the data point for monkey N416 was included within the cluster for the aged group (Fig. 5C).
sequencer (NGS). Specifically, 21 candidates of progeroid-related genes were sequenced in 100 monkeys including monkey N416 (see Table S2). The results showed that two genes (NBN and DDB2) have N416-specific mutations out of the 100 monkeys. However, these mutations are synonymous substitutions in NBN genes and downstream mutations in DDB2 genes, respectively, suggesting that an impact of such mutations on phenotypic functions is thought to be very weak.

Discussion

In our institute, we found a child Japanese monkey (monkey N416) who exhibited deep skin wrinkles and bilateral cataract within one year after birth. The following abnormalities were also found in this monkey: retardation of fibroblast proliferation at passage 5, appearance of senescent cells, deficiency in DNA repair, impaired glucose metabolism, altered hyaluronan metabolism, poikiloderma, shrinkage of the cerebral cortex and the hippocampus, and decrease in the conduction velocity of peripheral sensory nerves. However, the growth of this monkey was not retarded.
before she was 3 years old. These findings suggest that monkey N416 may probably suffer from a certain type of progeria.

Cataract

Bilateral cataract in monkey N416 seemed severe. In an individual cage, this monkey frequently smelled food pellets. As normal monkeys do not smell ordinary food so frequently, such a behavior may result from weak vision caused by cataract.

Retardation of fibroblast proliferation

Retardation of fibroblast proliferation is characteristic of WS and HGPS [6,7]. Our cell-culture examination showed a prolonged duration for fibroblast proliferation at passage 5 in monkey N416 as compared to a normal control, at a single cell level. This suggests that this monkey is likely progeroid. At passages 8, on the other hand, the population growth activity in fibroblast cells derived from monkey N416 was as high as that from the infant monkey, and higher than that from the aged monkey. This discrepancy can be accounted for by postulating that the difference in the manner of fibroblast proliferation in these experiments may be attributed to the difference in the passage number: passage 5 cells used for single-cell level analysis vs. passage 8 cells used for population level analysis. To address this issue, further investigations, for example flow cytometry of cell cycle constitution in early and late cultures, are needed.

Appearance of senescent cells

Acidic β-galactosidase is known to be a good hallmark of senescent cells [8]. Fibroblasts from HGPS patients express strong β-galactosidase activity [9]. Though it was previously reported that oxydative stress did not increase senescence-associated β-galactosidase activity in fibroblasts with WS [10], knockdown of RECQ2, RECQ3, and RECQL4 resulted in an increase in β-galactosidase-positive fibroblasts [11]. Our cytochemical analysis showed that the ratio of β-galactosidase-positive fibroblasts in monkey N416 was lower than in the aged monkey, and higher than in the infant

Figure 4. Shrinkage of the cerebral cortex and the hippocampus in monkey N416 was revealed with MRI. A,C,D: Parasagittal images of monkey N416 (1 year and 5 months old), an age-matched control, and an aged monkey (28 years old), respectively. Note that expansion of the cerebral sulci (arrows) in monkey N416 is as large as and expansion of the lateral ventricles (arrowheads) is larger than those in the aged monkey. This is probably ascribed to shrinkage of the cerebral cortex and the hippocampus. B: Coronal image of monkey N416. Seen is prominent expansion of the lateral ventricles (arrowheads).

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Figure 5. Conduction velocity of peripheral sensory but not motor nerve was slower in monkey N416. Measured from the ulnar nerve. A: Sensory conduction velocity (SCV). The SCV in monkey N416 and aged monkeys was slower than that in adult and child monkeys. **: p<0.01, *: p<0.05 (one-way ANOVA and Bonferroni post
monkey. Considering the difference in the number of passage of fibroblasts (N416 and aged 5, infant 7) and the age (infant 0 year, N416 3 years, aged 21 years), the ratio of senescent fibroblast cells in monkey N416 could be reasonable as a result of aging.

Deficiency in DNA repair
DNA repair deficiency is a common feature of HGPS [12] and other progeroid syndromes. RECQ3, RECQL4, and RECQ2 encode helicases, involved in DNA repair [1]. Our AP site assay have revealed that DNA repair does not simply deteriorate with the age, and that clear deficiency in DNA repair is observed in monkey N416. These findings strongly support that monkey N416 may suffer from a type of progeria.

Impaired glucose metabolism
It is well known that diabetes mellitus (DM) develops in most patients with WS [13]. Thus, we examined the stable DM marker, HbA1c. Levels of HbA1c in aged, adult, and child monkeys were as high as that in healthy rhesus (<4.7) [14] and long-tailed (4.4±0.1) [15] monkeys. However, HbA1c in monkey N416 was 4.8, estimated as “glucose impaired”, and significantly higher than in adult monkeys. This suggests that monkey N416 exhibits glucose metabolism impairment in accordance with WS patients, though she has not developed DM yet.

Altered hyaluronan metabolism
Hyaluronan is a key molecule to shape skin characteristics. For example, the Shar-pei dog and human patients with similar abnormal deep skin wrinkles share impairment in hyaluronan metabolism. Skin wrinkles in the Shar-pei dog results from upregulation of hyaluronic acid synthase 2 [16]. Deep skin wrinkles seen in human patients are correlated with an elevated hyaluronan concentration in serum [17]. Also, the hyaluronan level in WS is relatively high in urine [18]. The urinary hyaluronan level in monkey N416 was significantly higher than in child monkeys and was located between human WS patients and healthy controls [19]. The urinary hyaluronan level increased with the age in Japanese monkeys like humans, though the level itself was lower [18].

Poikiloderma and other chronic pathological changes
Based on the present pathological findings on the skin (slight acanthosis in epidermis, atrophy in dermis, and normal feature in subcutis, angiectasia in the superficial vascular plexus of dermis, and slight cellular infiltration of monocellular cell and eosinophilis in the superficial perivascular region of dermis), it can be considered that monkey N416 has poikiloderma with superficial telangiectasia. As poikiloderma is a representative feature of RTS, and BS [1,19], this favors that such pathological features may be ascribable to progeria.

In addition, there were chronic pathological changes in monkey N416, such as slight cardiac hypertrophy and renal pyelonephritis. It should be emphasized here, however, that these are quite rare in normal child monkeys.

Nervous systems
Expansion of the cerebral sulci and lateral ventricles, due to shrinkage of the cerebral cortex and the hippocampus, is a prominent aging-related change in the central nervous system of humans and rhesus monkeys [20]. In the present study, our MRI analysis indicated that the same event occurred in monkey N416 as well as in an aged control, but not in a child control. It remains to be known whether there are any histological changes (i.e., neuronal degeneration/loss) or molecular events that may be caused by aging.

The conduction velocity of peripheral sensory and motor nerves decreases when demyelination progresses [21]. In our experiments, we analyzed the conduction velocity of the ulnar nerve in monkey N416 by comparing with those in aged, adult, and child monkeys. The sensory conduction velocity (SCV) was significantly lower in monkey N416 and the aged monkey group than in the adult and child monkey groups. The SCV in adult and child monkeys was similar to that reported in the baboon [22]. Thus, it is most likely that demyelination of the sensory nerve may occur not only in the aged monkeys, but also in monkey N416. By contrast, the motor conduction velocity (MCV) in monkey N416 was not different from that in aged, adult, or child monkeys. The MCV measured in our study was similar to that reported in rhesus and long-tailed monkeys [23,24]. Further investigations are needed to clarify whether demyelination of the sensory nerve may occur in monkey N416 and whether the discrepancy between the SCV and the MCV may be ascribable to demyelination.

Progeroid-related genes
No individual-unique mutations of known progeroid-related genes were found in monkey N416. Some point mutations with reference to rhesus sequences were shared with normal monkeys. Therefore, it can be considered that phenotypes of monkey N416 are not simply attributed to such point mutations. To survey candidate genes responsible for progeroid phenotypes observed in monkey N416, further intensive analyses of the whole genome, at least at the exome level, are indispensable.

Comparison with major progeroid syndromes
As described above, monkey N416 displayed eminent symptoms of premature aging, such as bilateral cataract, retardation of fibroblast proliferation, appearance of senescent cells, deficiency in DNA repair, impaired glucose metabolism, altered hyaluronan metabolism, and poikiloderma. However, we could not detect any individual-unique mutations of known genes responsible for major progeroid syndromes. Typical facial features of HGPS include a small jaw, loss of hair, and scalp veins [2], none of which were observed in monkey N416. Moreover, this monkey did not show growth impairment, sclerotic skin, or joint contractures, each of which is characteristic of HGPS [25]. RTS is characterized by juvenile cataract, skeletal abnormality, proportionate short stature, and poikiloderma [1]. Juvenile cataract and poikiloderma were the common feature in both monkey N416 and RTS. WS is the most popular progeroid syndrome, though its frequency is less than 1/100,000 [26]. Some symptoms in monkey N416 were consistent with those in WS, including bilateral cataract, increased urinary hyaluronan, increased HbA1c, and retarded cell proliferation [18]. However, many symptoms seen in WS emerge only after adolescence [13], whereas several symptoms appear as early as 1 year old in monkey N416. In addition, there was no sign of osteoporosis, ulceration or atrophy of the skin, or calcification of the Achilles tendon [27], suggesting that monkey N416 does not suffer from WS. Progeroid phenotypes manifested in monkey
N416 are similar to in some aspects, but rather different from those in such major progeroid syndromes in humans (see Table 1).

In conclusion, the present study is the first to report a naturally occurring primate model with sporadic symptoms of premature aging. Further investigations especially with genome analyses and cell cultures will help us understand normal aging as well as progeria in primates including humans.

Supporting Information

Table S1  Sequence of primers used in PCR.

(XLS)

Table S2  Candidates for progeroid-related genes. There were N416 specific mutations in two progeria-related genes (NBN and DDB2). In the NBN, heterozygous synonymous mutation was found at exon 4, positioning of chr8:92,628,839 (coordinated by rheMac2), and in the DDB2, homozygous mutation was found at the downstream region, positioning of chr14:24,879,090 (also coordinated by rheMac2).

(XLS)

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Table S3  Levels of biomarkers in serum and urine, which showed no difference among N416 and age groups.

(XLS)

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Author Contributions

Conceived and designed the experiments: TO HI YG HH MT. Performed the experiments: TO HI YG HH. Analyzed the data: TO HI YG MI HH. Contributed reagents/materials/analysis tools: TO HI YG MI HH. Wrote the paper: TO HI YG HH MT.