Proof of Gene Doping in a Mouse Model with a Human Erythropoietin Gene Transferred Using an Adenoviral Vector

Takehito Sugawara 1,†, Takuro Nakano 2,†, Shin-ichiro Fujita 1,†, Yuki Matsumoto 3, Genki Ishihara 3, Kai Aoki 1,4, Koki Yanazawa 2, Seiko Ono 2, Shinsuke Tami 1, Lev Manevich 6,7, Haruna Ueda 8, Noriyoshi Ishibashi 9, Kensihiro Tamai 9, Yasuharu Kanki 1, Yasuko Yoshida 1,10, Koichi Watanabe 11, Tohru Takemasa 11, Yasushi Kawakami 3, and Kazuhiro Takekoshi 1*

1 Laboratory of Laboratory/Sports Medicine, Division of Clinical Medicine, Faculty of Medicine, University of Tsukuba, 1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan; take0716@krf.biglobe.ne.jp (T.S.); shin.ichiro.fujita.03@gmail.com (S.-i.F.); K-Aokitsuku@md.tsukuba.ac.jp (K.A.); yasuukk1220@gmail.com (Yasuharu Kanki); y-yoshida@ttius.ac.jp (Y.Y.); y-kawa@md.tsukuba.ac.jp (Yasushi Kawakami); k-takemd@md.tsukuba.ac.jp (Kazuhiro Takekoshi)
2 Master’s Program in Medical Sciences, Graduate School of Comprehensive Human Sciences, University of Tsukuba, 1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan; s2021408@s.tsukuba.ac.jp (T.N.); s1921312@s.tsukuba.ac.jp (K.Y.); jimbemilkybonbon330@gmail.com (S.O.)
3 Research and Development Section, Anicom Specialty Medical Institute Inc., 2-6-3 Chojamachi 5F, Yokohamashi-Nakacho, 231-0033, Kanagawa, Japan; ymatsumoto.ac@gmail.com (Y.M.); genki.ishihara@ani-com.com (G.I.)
4 Japan Society for the Promotion of Science, Kojimachi Business Center Building, Kojimachi, Chiyoda-ku, Tokyo 102-0083, Japan
5 Doctoral Program in Sports Medicine, Graduate School of Comprehensive Human Sciences, University of Tsukuba, 1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan; tama1994@outlook.jp (S.T.)
6 Doctoral Program in Biomedical Sciences, Graduate School of Comprehensive Human Sciences, University of Tsukuba, 1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan; lew.manewitsch@gmail.com (L.M.)
7 Department of Experimental Pathology, Faculty of Medicine, University of Tsukuba, 1-1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan
8 Analyst-Accenture Technology, Intelligent Platform Services, Accenture Japan Ltd, Akasaka Intercity AIR 1-8-1 Akasaka, Minato-ku, Tokyo 107-8672, Japan; hanamoo1995@gmail.com (H.U.)
9 Tsukuba i-Laboratory LLP, 2-1-17 Amakubo, Tsukuba 305-0005, Ibaraki, Japan; ishibashi@tsukuba-i-lab.com (N.I.); tamai@tsukuba-i-lab.com (Kenshiro Tamai)
10 Department of Medical Technology, Faculty of Health Sciences, Tsukuba International University, 6-20-1 Manabe, Tsuchiura, Ibaraki 300-0051, Japan
11 Faculty of Health and Sport Sciences, University of Tsukuba, 1-1-1 Tennodai, Tsukuba 305-8577, Ibaraki, Japan; watanabe.koichi.ga@u.tsukuba.ac.jp (K.W.); takemasa.tohru.gm@u.tsukuba.ac.jp (T.T.)
* Correspondence: k-takemd@md.tsukuba.ac.jp; Tel.: +81-29-853-3209
† These authors contributed equally to this work.

Abstract: The World Anti-Doping Agency (WADA) has prohibited gene doping in the context of progress in gene therapy. In addition, there is a risk of the EPO gene being applied in gene doping among athletes. Along with this, development of a gene-doping test has been underway worldwide. Here, we had two purposes: to develop a robust gene doping mouse model using the human EPO gene (hEPO) transferred using recombinant adenovirus (rAdV) as a vector and to develop a detection method to prove gene doping using this model. The rAdV including the hEPO gene were injected intravenously to transfer the gene to the liver. After injection, the mice developed significantly increased red blood cell counts in whole blood and increased gene expressions of hematopoietic markers in the spleen, indicating successful development of the gene doping model. Next, we detected direct and indirect proof of gene doping in whole blood DNA and RNA using qPCR assay and RNA sequencing. Proof was detected in one drop of whole blood DNA and RNA over a long period; furthermore, the overall RNA expression profiles significantly changed. Therefore, we have advanced detection of hEPO gene doping in humans.

Keywords: Gene doping; Gene therapy; Erythropoietin; Adenoviral vector; Sports; Athlete; RNA sequencing
Graphical abstract
1. Introduction

Doping is the act of using prohibited substances and/or methods in sports to enhance athletic performance and success by improving physical performance [1]. The World Anti-Doping Agency (WADA), which was established in 1999, is involved in scientific research on doping, anti-doping education, development of anti-doping strategies, and monitoring of the World Anti-Doping Code to ensure soundness and fairness in sports worldwide [2]. However, despite WADA’s substantial efforts, doping has not been eradicated from competitive sports.

The “International Standard Prohibited List” [3] stipulated in the “World Anti-Doping Code 2021” [4], which is published with annual revisions by the WADA, describes various formulations and methods used in doping. This includes “gene doping” as abuse of gene therapy technology. Therefore, our research group has been conducting research for more than five years to establish an examination method for gene doping [5-7].

Gene therapy has rapidly evolved as a new treatment method for single hereditary diseases and acquired diseases, and these treatments have yielded many beneficial results for human diseases [8]. In fact, gene therapeutics, as represented by Collategen (AnGes, Ibaraki, Osaka, Japan), Zolgensma (Novartis Gene Therapies, Chicago, IL, USA), and YESCARTA (Kite Pharma, Santa Monica, CA, USA), have been approved for treatment of human disease. Recently, gene therapy technology has also been used to develop vaccines based on mRNA [9] or adenoviral vectors [10], against coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2). For that reason, the WADA has raised concerns about the abuse of the gene therapy technology for doping. However, there is no robust and universal method to detect gene doping at present. On the other hand, the WADA has recently established laboratory guidelines for PCR methods upon gene doping tests. Even though, in the guidelines, basic validation methods in the PCR method are described in detail [11], there is no description of detection methods for individual target genes or vectors. Therefore, it would be desired to develop detailed detection methods that can detect any types of gene doping.

In particular, the erythropoietin (EPO) gene can be abused in gene doping because EPO and EPO-related formulations had history that it is frequently used in doping to enhance the endurance performance of athletes in cycling, boxing, athletics, and rowing [12-16]. Thus, EPO and EPO-related formulations are included on the “International Standard Prohibited List” [3] published by WADA. According to the “Wiley database on Gene Therapy Trials Worldwide” [17], recombinant adenovirus (rAdV) is the most frequently used vector in clinical trials for gene therapy. Thus, there is a strong possibility that combinations of the EPO gene and rAdV are being used for gene doping in human athletes. In modern times, establishing a method for detecting gene doping using this combination is urgently needed.

Although some previous studies have attempted to establish animal models of gene doping by using naked plasmid vectors including the EPO gene [6,18,19], they did not evaluate the phenotypic effects of EPO gene doping, such as changes in the number of red blood cells (RBCs), gene and protein expression of EPO, and secretion levels of EPO in blood. For these reasons, the models proposed in these studies may lack accuracy and robustness and be limited for developing detection methods for gene doping. In addition, no reports have established a gene doping model using the combination of the EPO gene and rAdV. Comprehensively considering these previous studies and history, we recognized that a robust model to develop a detection method for direct or indirect proof of gene doping using rAdV and the human EPO (hEPO) gene (rAdV-hEPO) was needed.

As direct proof, we considered that the detection of viral vector-specific DNA fragments by a TaqMan qPCR assay would be the most reasonable approach because the principle underlying this assay has also been used successfully in testing for SARS-CoV-2. In addition, as indirect proof, we considered that monitoring fluctuations of RNA expression in whole blood could be applied to the parameters of the Athlete Biological Passport...
(ABP) because its fundamental principle is to monitor selected biological variables that indirectly reveal the effects of doping over time rather than to attempt to detect the doping substance or method itself [20]. Furthermore, the WADA describes that the ABP can be used to identify athletes requiring further attention through intelligent, timely interpretation of passport data and can notably be used as a complement to analytical methods to further refine and strengthen overall anti-doping strategies [20]. In other words, the detection of indirect proof is also important to identify athletes requiring further attention and to support the findings as strongly as direct proof in gene doping tests. Therefore, in this study, total RNA sequencing (RNA-seq) using whole blood RNA from an animal model was also performed to identify RNAs as novel indirect proof based on the concept of the ABP.

In summary, the primary objectives of this study were the establishment of a robust gene doping mouse model using rAdV-hEPO, the discovery of RNA as indirect proof based on the concept of the ABP, and the development of detection methods of direct and indirect proof for gene doping.

2. Materials and Methods

2.1. Creation of the rAdV-hEPO vector

To establish a robust gene doping mouse model injected with rAdV-hEPO, a viral vector was created using the following procedures. Three plasmids, namely, (p) hEPO-Myc-DDK-tag (Cat#RC21077S1; ORIGENE, Rockville, MD, USA), pENTR4 (Cat#A10465; Thermo Fisher Scientific, Waltham, MA, USA), and pAd/CMV/V5-DEST (Cat# V49320; Thermo Fisher Scientific; can make rAdV type 5), were used to create the rAdV-hEPO vector. First, PCR was performed to amplify the hEPO gene including the myc and DDK (FLAG) tags and the restriction enzyme sites of EcoRI/Not I from the phEPO-Myc-DDK-tag using PCR enzymes (KOD-Plus-, Cat# KOD-201; TOYOBO, Osaka, Osaka, Japan). The PCR amplicon was subjected to gel electrophoresis, after which the amplicon DNA was extracted and purified from the gel. After digestion of the purified amplicon and pENTR4, a homologous recombination reaction was induced in the mixture using an In-Fusion HD Cloning Kit (Cat# 63964; Takara Bio, Kusatsu, Shiga, Japan). The obtained plasmid (pENTR4-hEPO) was mixed with pAd/CMV/V5-DEST, and an LR recombination reaction was induced using Gateway LR Clonase II Enzyme mix (Cat# 11791020; Thermo Fisher Scientific). Finally, the obtained plasmid (pAd/CMV/V5-DEST-hEPO) was digested using a Pac1 restriction enzyme (Cat# R0547; New England Biolabs, Ipswich, MA, USA) and transfected into HEK 293A cells (Cat#R70507; Thermo Fisher Scientific) with a transfection reagent (PEI MAX, Cat#24765-1; Polysciences, Warrington, PA, USA) to amplify rAdV type 5 including the hEPO gene cassette. The amplified rAdV-hEPO vectors were purified by CsCl density-gradient ultracentrifugation followed by gel filtration with a PD-10 Column (Cat#17085101; cytiva, Marlborough, MA, USA) according to the protocol described by Takeuchi et al. [21,22]. The viral particles (VPs) were suspended in 10% glycerol/phosphate-buffered saline (PBS), and the concentration of rAdV VPs was measured using a spectrophotometer according to the method described by Sweeney and Hennessey [23].

2.2. Animal experiments

All animal experiments in this study were approved by the Animal Care Committee, University of Tsukuba (approval numbers: 20-361). Seven-week-old male ICR mice were purchased from CREA Japan (Meguro, Tokyo, Japan) and then subjected to a 1-week acclimation period. The mice were bred and maintained in an air-conditioned animal house under specific pathogen-free conditions and subjected to a 12/12-h light/dark cycle. The mice were fed standard mice pellets and water ad libitum. At the start of the experiments,
the age of the mice was 8 weeks. The animal experiments were broadly divided into short- and long-term experiments.

The short-term experiments were conducted to establish a gene doping mouse model with rAdV-hEPO and to develop methods to detect direct and indirect proof of gene doping. An overview of these experiments is shown at Fig. 1A. After 1 week of acclimatization, mice were randomly assigned to the Control (Con.) or rAdV-hEPO groups. The mice in the rAdV-hEPO group (n = 16; named AdEPO mice) received injections of the AdV-hEPO vector (4.0 × 10^11vp/100 µL/mouse) into the orbital sinus under systemic isoflurane inhalation anesthesia. The mice of the Con. group (n = 12) received injections of the 10% glycerol/PBS (100 µL/mouse) buffer used to suspend the rAdV. Five days after the injection, whole blood was obtained with EDTA-2Na as an anticoagulant from the inferior vena cava under systemic isofurane inhalation anesthesia, after which the mice were euthanized. The collected whole blood was subjected to preprocessing for further analysis. Samples of liver and spleen were also harvested and flash-frozen in liquid nitrogen until further analysis. Other samples of spleen tissues were immersed into 10% formalin neutral buffer solution overnight to obtain paraffin block specimens. Supplementary experiments were conducted using the rAdV-ZsGreen1 (ZSG) vector to confirm the effects of hEPO-free rAdV on the phenotypes.

Long-term experiments were conducted to investigate how long direct and indirect proof of gene doping could be positively detected from approximately one drop of whole blood. The overview of these experiments is shown in Fig. 5A. After 1 week of acclimatization, small whole blood samples (approximately 100 µl; 2 drops) from mice (n = 12) were collected with EDTA-2Na using cuts made approximately 2 mm from the tail tip (Pre-time point). Then, the AdV-hEPO vector was injected using the methods described above. Subsequently, blood samples were continually collected until 30 days after injection using the same methods. The collected blood samples were divided into two samples of approximately 50 µL each, and DNA and RNA were extracted from these samples and subjected to further analyses.

2.3. Measurements of general hematopoietic markers

Hematological indicators of RBCs, namely, the RBC count, hemoglobin (Hgb level, and hematocrit (HCT) value, from the whole blood samples obtained in the short-term experiments were measured on an automatic blood analyzer (Celltac α MEK6458; NICHON KODEN, Shinjuku, Tokyo, Japan) using 50 µL of whole blood.

2.4. Western blotting

To confirm the expression of hEPO-protein in the liver in the short-term experiment, western blotting (WB) was performed. To extract the total protein from liver, liver specimens were homogenized in lysis buffer (50 mM Tris-HCl pH 7.4, 150 mM NaCl, 1% NP40, 1 mM EDTA) with a protease inhibitor cocktail (Cat# 25955-11; Nacalai Tesque, Nakagyo, Kyoto, Japan) using a bead crusher (TissueLyser LT, Cat#85600; QIAGEN, Hilden, NRW, Germany). The homogenized liver lysate was then centrifuged at 12,000 × g for 15 min, and the supernatant was collected. The concentration of each protein was measured using a BCA kit (Cat#T9300A; Takara Bio) and adjusted to 2 µg/mL. The supernatant was mixed with 2× loading buffer containing 2-mercaptoethanol and denatured at 95°C for 5 min. Subsequently, 10-µL samples including 10 µg of protein were subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) using 15% gel at 140 V for 70 min, after which the separated proteins in the gel were transferred to a polyvinylidene fluoride (PVDF) membrane using the wet transfer method at 40 V overnight. This membrane was blocked with TBS-T buffer (50 mM Tris-HCl, 150 mM NaCl, 0.05% Tween-20) including 5% skim milk for 120 min and then washed with TBS-T buffer 3 times for 10 min. The membrane was incubated with the primary antibody for EPO (Cat#sc-80995; Santa Cruz Biotechnology, Dallas, TX, USA) at 100-fold overnight at 4°C with gentle shaking and then washed with TBS-T buffer three
times for 10 min. The washed membrane was incubated with anti-rabbit IgG HRP-linked Antibody (Cat#7074, 3,000-fold; Cell Signaling Technology, Danvers, MA, USA) with gentle shaking for 30 min at room temperature. After washing the membranes three times for 10 min, the target protein bands were visualized with a chemiluminescence reagent (EzWestBlue; Cat#WSE-7140; ATTO, Taito, Tokyo, Japan), on ImageQuant LAS 4000 (Cytiva). The band images were exported as 16-bit TIFF images. The luminance of the bands of the TIFF images were quantified using ImageJ Fiji (ver. Java 8). The antibody for glyceraldehyde-3-phosphate dehydrogenase (GAPDH, Cat# 60004-1-lg, 3,000-fold; Proteintech, Rosemont, IL, USA) was also used as the loading control with anti-mouse IgG HRP-linked antibody (Cat#7076, 3,000-fold; Cell Signaling Technology).

2.5. Enzyme-linked immunosorbent assay

Enzyme-linked immunosorbent assay (ELISA) was performed to confirm secretion of hEPO into the blood from the liver in AdEPO mice in the short-term experiment. Anti-hEPO (Cat#500-P318; PeproTech, Cranbury, NJ, USA) was diluted to 0.5 µg/mL in PBS as a capture antibody, and 100 µL of the antibody solution was then applied into an ELISA plate as duplicate measurements. The plate was incubated at 4°C overnight and then washed with PBS containing 0.05% Tween 20 (PBS-T) four times; then, 100 µL of the plasma samples diluted 10-fold with PBS were added to the wells and incubated for 1 h at room temperature. Solutions of hEPO recombinant protein (Cat#100-64; PeproTech) were also added to generate a standard curve between 100 ng/mL and 10 pg/mL of PBS. After incubation, the plate was washed in the same way as described previously, and a 100-µL solution of a detection antibody of biotinylated anti-hEPO (Cat#500-P318BT; PeproTech) diluted with PBS-T was applied to the wells at a concentration of 0.25 ng/µL, followed by incubation for 1 h at room temperature. After incubation, the plate was washed in the same way as described previously, and a 100-µL solution of HRP-conjugated streptavidin (Cat# SA00001-0; Proteintech) diluted 5,000-fold with PBS-T was applied to the wells. The plate was incubated at room temperature for 30 min and then washed in the same way as described previously. After washing, a 100-µL mixture of coloring reagent and substrate (ELISA POD Substrate A.B.T.S Kit; Cat#14351-80; Nacalai Tesque) was applied and incubated for 20 min, followed by application of 100 µL of stop reagent for the coloring reaction. Finally, the absorbance at 405 nm with a reference at 600 nm was measured using a microplate reader. Using the absorbance data, a standard curve as a 4-parameter logistic of the Rodbard was created in ImageJ Fiji (Life-Line version, updated on 2017 May 30), and the concentration of hEPO in the plasma was calculated based on the standard curve ($R^2$=0.99) as duplicate measurements.

2.6. TB Green qPCR assay for tissue RNAs

To confirm the expression of hematopoietic marker genes in the liver and spleen in the short-term experiment, the TB Green qPCR assay was performed as an intercalation method. RNA extraction from the liver and spleen was performed using RNAiso Plus (Cat#9180; Takara Bio) according to the manufacturer’s instructions. The extracted RNA solution in Milli-Q Water (Merck Millipore, Burlington, MA, USA) was diluted and adjusted to a concentration of 100 ng/µL. Then, 500 ng of RNA was used to prepare cDNAs using PrimeScript RT Master Mix (Cat#RR036A; Takara Bio) according to the manufacturer’s instructions. The cDNAs were diluted ×10 using Milli-Q Water and subjected to a quantitative real-time PCR (qPCR) assay based on the intercalator-fluorescence dye. The qPCR assay was performed to quantify the gene expression of hematopoietic markers in the liver and spleen using TB Green Premix Ex Taq II (Cat# RR820; Takara Bio) with primers from QuantStudio 5 Real-Time PCR Systems (Thermo Fisher Scientific) as duplicate measurements. The targeted gene list and primer sequences are shown in Supplementary Table S1. The template volume and primer concentration were 2 µL and 100 nM, respectively, for a total reaction volume of 10 µL per well. Negative control wells
were also established using pure water instead of the template. The conditions for thermal cycling were 95°C for 5 min, followed by 40 cycles of 95°C for 2 s and 60°C for 20 s and a melt curve stage. Subsequently, the \( \Delta \Delta^C T \) method referencing 18s rRNA was used to calculate the relative gene expression values.

2.7. Immunohistochemistry

Immunohistochemistry (IHC) was performed to confirm the expression and localization of GATA1, a common hematopoietic marker, in the short-term experiment. In this experiment, 4-μm sections of spleen on slides were deparaffinized with Lemosol A (Cat# 126-04413; FUJIFILM Wako Pure Chemical Corporation, Osaka, Osaka, Japan) and rehydrated using ethanol and running water. After 3 min of rinsing with running water, the endogenous peroxidase in the tissues was removed by treatment with 3% hydrogen peroxide/methanol for 10 min. After washing with PBS for 1 min with gentle shaking, the sections were placed in 0.01 M citrate buffer and subjected to antigen activation at 121°C for 10 min. Then, the sections were gently washed with PBS and blocked with 5% bovine serum albumin (BSA)/PBS-T for 1 h at room temperature. The primary antibody for GATA1 (Cat#10917-AP; Proteintech) diluted 100-fold with 1%BSA/PBS-T was applied to the sections and incubated at 4°C overnight. On the next day, after washing the slides with PBS-T three times for 15 min, anti-rabbit IgG HRP-linked antibody (Cat#7074; Cell Signaling Technology) diluted 1:100 with 1% BSA/PBS-T was applied to the sections and incubated for 1 h at room temperature after washing with PBS. Subsequently, 3,3-diaminobenzidine tetrahydrochloride (DAB) solution in the peroxidase stain DAB Kit (Cat#2598-50; Nacalai Tesque) was applied to the section, and the section was incubated for 5 min at room temperature, followed by washing with PBS and counterstaining with hematoxylin. The sections were then dehydrated and mounted using Malinol (Cat#2009-3; MUTO PURE CHEMICALS, Bunkyo, Tokyo, Japan) and observed under a microscope (BZ-X710; Keyence, Osaka, Osaka, Japan).

2.8. DNA extraction from whole blood samples

A phenol/chloroform/isoamyl alcohol solution (Cat#25970-56; Nacalai Tesque) was used to extract total DNA from 100 μL (short-term experiment) or 50 μL (long-term experiment) of whole blood in accordance with the manufacturer’s instructions. The pellets of the DNA were dissolved in 50 μL of Milli-Q Water, and the DNA solutions were subjected to the TaqMan qPCR assay to detect direct proof of gene doping.

2.9. TaqMan qPCR assay

Our previous studies have shown that whole blood DNA contains high levels of viral genome DNA fragments as direct proof [5-7]; therefore, we performed the TaqMan qPCR assay with whole blood DNA. For the TaqMan qPCR assays to detect direct proof of gene doping in whole blood DNA in the short- and long-term experiments, the primers and TaqMan probes were designed to target the EPO gene (2 types), TkpA, Hexon, and CMVp to ensure specific amplification of the rAdV-hEPO genome using Primer-BLAST (NIH National Library of Medicine, Bethesda, MD, USA). The primers and TaqMan probes for the 2 types of hEPO genes were prepared with exon-exon junctions, which is a nonamplifying form in the human genome [6,24,25]. An overview of these design strategies is shown in Fig. 2-A. The primers and probes were also checked for specificity with in silico PCR using Primer-BLAST; these evaluations confirmed that there was no amplification from the human and mouse genomes. The sequences of the primers and TaqMan probes are shown in Supplementary Table S1. The primers and probes in a double quencher system were systemized by Integrated DNA Technologies (Coralville IA, USA). Next, the TaqMan qPCR assay was performed in duplicate to detect direct proof in whole blood DNA by absolute quantification using PrimeTime Gene Expression Master Mix (Cat# 1055771; Integrated DNA Technologies) with the primers and TaqMan probes on QuantStudio 5 Real-Time PCR Systems (Thermo Fisher Scientific). The tem-
plate DNA volume was 2 µL, and the primer and probe concentrations were 200 nM and 100 nM, respectively, for a total reaction volume of 10 µL per well. Negative control wells were also established using pure water instead of a template. phEPO-Myc-DDK-tags including the hEPO gene or pAd/CMV/V5-DEST including TkpA, Hexon and CMVp were used at 100 pg/µL to prepare a standard curve for absolute quantification, and the range of the standard curve was set to $1.19 \times 10^7$ to 1.9 copies/µL for the phEPO-Myc-DDK-tag and $2.53 \times 10^6$ to 10.1 copies/µL for the pAd/CMV/V5-DEST. The conditions of thermal cycling for all primer-probe pairs were 95°C for 5 min, followed by 40 cycles of 95°C for 2 s and 60°C for 20 s. All standard curves had $R^2 > 0.98$.

2.10. Sanger sequencing

After the TaqMan qPCR, solutions including the amplicon were pooled in a 1.5-mL microtube and then subjected to electrophoresis using 2% agarose gel. The bands of the DNA amplicons were visualized, cut, and purified using a NucleoSpin Gel and PCR Clean-up kit (Cat# 740609; Takara Bio). Next, 5 ng of the purified DNA was subjected to Sanger sequencing via outsourcing to an external company (GENEWIZ, Shinagawa, Tokyo, Japan) to check the sequence of the DNA amplicons. The Sanger sequencing data were analyzed with CLC Sequence Viewer ver. 8.0 (QIAGEN) and BioEdit ver. 7.2.5 (developer: Tom Hall).

2.11. Total RNA-seq

Total RNA-seq was performed to identify gene as novel indirect proof based on concept of the ABP. Total RNA of the mice in the short-term experiments was extracted from 100 µL of whole blood using RNeasy Blood (Cat#9112; Takara Bio) according to the manufacturer’s instructions. The RNA pellets were dissolved in 30 µL of Milli-Q Water, and the RNA solutions of eight samples (Con.: N = 4 and rAdV-hEPO: N = 4) were checked for integrity using Agilent RNA 600 Nano Kit (Cat# 5067-1511; Agilent Technologies, Santa Clara, CA, USA) on the Bioanalyzer (Agilent Technologies). The RNA Integrity Number (RIN) of all samples was 8.8 or higher; thus, the RNAs of all eight samples could be subjected to library preparations for total RNA-seq. Using 300 ng of the RNAs from each sample, libraries were created using NEBNext Ultra II RNA Library Prep Kit for Illumina and NEBNext rRNA Depletion Kit v2 (Cat# E7770S and E7400L; New England Biolabs) according to the manufacturer's instructions, and the final PCR cycle was 12. Concentrations and size distributions of the libraries were measured using an Agilent DNA 7500 kit (Cat#5067-1506; Agilent Technologies) with Bioanalyzer. All samples were passed for analyses on NGS equipment.

The libraries were pooled, and the concentrations were adjusted to 1 nM. The pooled libraries were subjected to denaturation and neutralization. Subsequently, the libraries were diluted to 1.8 pM and then applied for an NGS run using NextSeq500/550 v2.5 (75 Cycles) Kits (Cat#20024906; Illumina, San Diego, CA, USA) in the NextSeq 500 System (Illumina). The sequencing was performed with paired-end reads of 36 bases. After the sequencing run, FASTQ files were exported, and the basic information of the NGS run data was checked on CLC Genomics Workbench 20.0.3 software (QIAGEN). As a quality score of the reads, a PHRED-score over 20 was confirmed for 99.73% of all reads, indicating the success of the run. The read number was approximately 37 to 45 million per sample as paired-end reads.

2.12. Bioinformatics analysis

The following analysis was performed to identify genes as novel indirect proof based on the concept of the ABP using NGS run data. FASTQ files were mapped to the mouse genome (mm 10) using the CLC Genomics Workbench software (QIAGEN). A statistical differential expression test was performed by empirical analysis using the Differential Expression in Two Groups tool in the software. Differentially expressed genes (DEGs) defined by the cutoff levels were considered with false discovery rate
(FDR) < 0.01 and a 3-fold change cutoff. A principal component analysis (PCA) plot was created using the CLC software. Transcripts Per Kilobase Million (TPM) was used as an expression value for figure visualizations. The low read count genes were filtered by 0 in all samples. Gene Set Enrichment Analysis (GSEA) software [26] was used to perform the enrichment analysis. Expression datasets and phenotype files were created and imported into GSEA software. According to "Using RNA-seq Datasets with GSEA", input normalized data were created with DEseq2 using TCC-GUI to perform GSEA [27]. The GSEA process was performed as the default setting with the h.all.v7.4.symbols, C2.cp.v7.4.symbols, C2.cgp.v7.4.symbols, and C5.go.v7.4.symbols datasets. Cell-type specific analysis (CTSA) was performed using TissueEnrich R package with the default settings [28] with RNA single cell type data from The Human Protein Atlas (https://www.proteinatlas.org/) to calculate cell-type specific gene enrichment. The names of the DEGs were matched to the corresponding human gene names with the biomart R package for conducting CTSA [29]. Tissue-specific expression analysis (TSEA) was performed using the TissueEnrich web tool (https://tissueenrich.gdcb.iastate.edu/) [28] with the Mouse ENCODE Dataset to calculate tissue-specific gene enrichment. The Database for Annotation, Visualization and Integrated Discovery (DAVID version 6.8, LHRI) web tool was used to elucidate the enrichment terms with default settings [30].

2.13. TB Green qPCR assay for whole blood RNA

To confirm whether the identified genes as indirect proof of gene doping showed reproducibility for the total RNA-seq results, a TB Green qPCR assay was performed for all samples (Con.: n = 12, rAdV-hEPO: n = 15) in the short-term experiment. RNA extraction from 100-µL whole blood samples was performed using the same methods mentioned in the “Total RNA-seq” section. Subsequently, the TB Green qPCR assay for the identified genes as indirect proof was performed using the same method described in the “TB Green qPCR assay for tissue RNAs” section. The primer sequences used in this section are shown in Supplementary Table S1.

In addition, to investigate the period for which the identified indirect proof could be positively detected, the same qPCR assay was also performed using 50-µL whole blood samples (n = 12) from the long-term experiment. In this experiment, cDNA concentrations were additionally measured using the QuantiFluor ssDNA System (Cat# E3190; Promega, Madison, WI, USA) according to the manufacturer’s instructions. Subsequently, the expression values of each targeted gene were normalized by their cDNA concentrations because the expression levels of the normalizer gene (18s rRNA) significantly changed according to the passage of days.

2.14. Statistical analysis

All data except the total RNA sequence data were statistically analyzed using GraphPad Prism version 9.0.2 (GraphPad, San Diego, CA, USA). All experimental data were first evaluated with the Shapiro–Wilk normality test to check the normality of the distributions. Subsequently, nonparametric tests were used for all data. For comparisons of 2 groups, the Mann–Whitney U test was performed. Comparisons of 3 or more groups were performed with Kruskal–Wallis H tests (one-way ANOVA of ranks) followed by a two-stage Benjamini, Krieger, and Yekutieli FDR procedure as a post-hoc test. A p value less than 0.05 was considered to indicate statistical significance. All graphs without data from the bioinformatics analysis of the total RNA-seq are shown as individual plots and medians with interquartile ranges.

3. Results

3.1. Mouse model of gene doping established using rAdV-hEPO
Fig. 1A shows an overview of this experiment. A phenotypic analysis performed 5 days after the injection of rAdV-hEPO showed a significant increase in the body weight of AdEPO mice compared with the control mice. The livers and spleens of AdEPO mice also showed significant hypertrophy (Fig. 1B-D). The RBC counts, Hgb levels, and HCT values, which served as hematological markers in whole blood, were also significantly increased (Fig. 1E-H), indicating the occurrence of hematopoiesis.

To confirm whether hEPO was expressed and secreted from the liver, qPCR, WB, and ELISA were performed using liver RNA, protein lysate, and plasma. The results showed that the total EPO protein expression was significantly elevated in AdEPO mice, and the hEPO hormone was detected in the plasma only in AdEPO mice along with positive expression of the hEPO gene in the liver (Fig. 1H-L). The raw data from the WB analysis are shown in Supplementary Figs. S1 and S2.

In addition, we confirmed whether the gene expression levels of the downstream targets of EPO in hematopoiesis signaling had changed in the livers and spleens of AdEPO mice. The results showed significantly upregulated expression levels of Gata1, Vegfa, and Vegfb in the livers of AdEPO mice. Moreover, the expression levels of Gata1, Trfr, and Gypa were drastically upregulated in the spleens of AdEPO mice (Fig. 1M). In the spleen, increased GATA1 protein expression was also confirmed using IHC. These results suggested that gene delivery to the liver, secretion of hEPO into blood, and hematopoiesis in the spleen were successfully achieved by injection of rAdV-hEPO, indicating the hematopoietic function of hEPO.

To investigate the effects of hEPO-free rAdV on these phenotypes and gene expression, we conducted a supplementary experiment using rAdV-ZSG as a form of hEPO-free rAdV that could produce green fluorescent protein. Induction of rAdV-ZSG did not yield phenotypes and gene expression related to hematopoiesis (Supplementary Fig. S3 A-I). Thus, rAdV alone had no hematopoietic effect in mice. Overall, these results clearly indicated the establishment of a gene doping model with rAdV-hEPO in this study.
Figure 1. Establishment of a gene doping model using rAdV-hEPO. The phenotypes and parameters related to hematopoiesis were assessed in the Con. (n = 12) and rAdV-hEPO (n = 15) groups. (A) an overview of the experiment, (B) a photograph of liver and spleen of a representative mouse, (C) body weight of the mice, (D) liver and spleen weight, (E) RBC count (10⁵/μL), (F) Hgb level (g/dL), (G) HCT (%), (H) expression of the hEPO gene in the liver, (I) representative images of total EPO and GAPDH on WB analysis of liver specimens, (J) quantification of the band intensity on WB, (K) a representative image of the actual reaction plate of ELISA for the hEPO protein, which was performed as duplicate measurements of 6 samples, (L) results of quantification with ELISA, (M) analysis of the expression levels of genes encoding hematopoietic markers in the liver and spleen, (N) micrographs of GATA1-stained spleen sections evaluated using IHC. N.D.: not detected. *p < 0.05, **p < 0.01, ***p < 0.001.

3.2. Direct proof of gene doping in whole blood DNA from AdEPO mice

Using specific primer-probe pairs (Fig. 2-A) and whole blood DNA in the TaqMan qPCR assay, the results provided direct proof of gene doping with statistical significance for any 5 primer-probe pairs (Fig. 2-B). The median copy numbers of the targeted DNA fragments were confirmed to be approximately 500-1,500 copies/μL in whole blood. However, positive detection was also confirmed in the control mice when using the primer-probe pairs to detect the Hexon region (Fig. 2-B; Pr-Hexon).

The DNA sequences of the amplicon for each TaqMan qPCR assay were confirmed using pooled amplicon samples with gel extraction using the Sanger sequencing method. The results showed correct sequences over 40 nucleotides in any primer-probe pair (Fig. 2C-G). Therefore, these results suggested that these primer-probe pairs can accurately detect direct proof in AdEPO mice as a gene doping model. On the other hand, it is possible that there were experimental contaminations of DNA fragments of the Hexon region.

Figure 2. Direct proof in whole blood DNA from AdEPO mice. (A) Strategy for designing specific primer-probe pairs for the viral genome. “Pr” indicates a primer–probe pair. (B) Detection of direct proof using the TaqMan qPCR assay performed with whole blood DNA from the control (n = 12) and AdEPO mice (n = 15). (C) to (G) Sanger sequence analysis of the pooled amplicon on the TaqMan qPCR assay using five primer probes; R.S. indicates a reference sequence that completely matches the amplicon over 40 nucleotides on any primer-probe pairs. N.D.: not detected. ***p < 0.001.

3.3. RNAs as novel-indirect proof of gene doping in whole blood RNA
Total RNA-seq and bioinformatics analysis was performed to identify potential indirect proof of gene doping using whole blood RNA. The PCA plot showed that the comprehensive gene expression profiles considerably differed between control and rAdV-hEPO mice (Fig. 3A). Gene expression profiles were relatively similar across the samples within each group, except for one control sample (Fig. 3A). The statistical analysis identified 1,128 DEGs (Fig. 3B). The DEGs included 94.95% protein-coding genes, 4.34% long non-coding RNA (lncRNA) genes, 0.35% small nucleolar RNA (snoRNA) genes, 0.27% small nuclear RNA (snRNA) genes, and 0.09% immunoglobulin constant germline (IGC) genes. The gene expression profile included 903 upregulated and 225 downregulated genes (Fig. 3C). Interestingly, GSEA predicted several gene sets associated with hematopoiesis-related terms (Fig. 3D, E, and F). In particular, the genes included in GATA1 target term are known to be related to RBC development and component functions (Fig. 3G). In addition, GSEA indicated that genes for specific elements of hemopoiesis-related mechanisms were significantly enriched, such as mitochondria-related electron transport chain, OXPHOS pathways, and metal ion homeostasis-related terms (Supplementary Fig. S4). Next, we hypothesized that DEGs might include RBC-specific genes that could be candidate genes for gene doping with rAdV-hEPO. Thus, CTSA was performed to identify potential cell type-specific genes in circulating RNA. Erythroid cell-specific genes were most significantly enriched on CTSA (Fig. 3H). In addition, DEGs were confirmed to have other blood cell type-specific gene changes although those cell types were not enriched on CTSA (Fig. 3I). Thus, these blood cell-specific genes, particularly RBC-specific genes, may serve as indirect proof of rAdV-hEPO gene doping (Fig. 3J).
Figure 3. Hematopoietic changes caused by rAdV-hEPO supported by whole blood RNA-seq. (A) PCA plot showing similarities between samples. (B) Bar plot and pie chart showing the 1,128 DEGs included in the total detected genes in whole blood and rates of the gene types. (C) Heat map indicating the DEG profile. Values in the rows are z-scores. GSEA showing enriched terms: heme metabolism (D), hematopoiesis mature cell (E), and GATA1 targets (F) terms. (G) Relative gene expressions levels (Con. = 1.0) showing gene expression in the GATA1 target term in Fig. 3F. Biological-related functions are described under gene names. The genes were visualized according to the GSEA enrichment results with a “Yes” value. Error bars indicate the standard error of the mean (SEM). (H) Bar plot showing CTSA to predict cell type-specific enrichment in DEGs. The gray line represents a significant threshold (p < 0.05). (I) Box plot showing single-cell gene normalized expression (“NX”) of DEGs in each cell type. Single-cell gene expression levels of DEGs were classified by cell type-specific definitions by TissueEnrich for each cell type. (J) Relative gene expression levels (Con. = 1.0) showing erythroid cell-enriched gene expressions in DEGs. Abbreviations: long non-coding RNA (lncRNA), small nucleolar RNA (snoRNA), small nuclear RNA (snRNA), and immunoglobulin constant germline gene (IGC gene).
Furthermore, to identify potential genes in whole blood that may reflect tissue status, we conducted TSEA, which showed that E14.5-liver-, bone marrow-, and spleen-categorized genes were enriched (Supplementary Fig. S5A). In addition, we noticed that the E14.5-liver-categorized genes overlapped with the bone marrow-categorized genes (Supplementary Fig. S5B and C). These 42 overlapping genes were enriched terms such as porphyrin-containing compound biosynthetic process, erythrocyte development, protoporphyrinogen IX biosynthetic process, heme biosynthetic process, and erythrocyte differentiation (Supplementary Fig. S5D). For example, Rhd, Kif1, Rhag, and Gata1 were included in the erythrocyte development term (Supplementary Fig. S5E). In particular, Gata1 was upregulated in whole blood (Supplementary Fig. S5F), consistent with the results for tissues (Fig. 1M). These results can suggest a hematopoietic status inside tissues after rAdV-hEPO injection, consistent with the tissue phenotype. In addition, the changes in the overall RNA expression profiles of these hematopoiesis-associated genes may serve as indirect proof of gene doping based on the concept of the ABP.

Next, we screened the single candidate genes that showed drastic changes in expression upon gene doping. These genes were identified by filtering high fold changes and high expression levels (Fig. 4A) because the gene should be measurable with high sensitivity on qPCR. Nine candidate genes were screened in this assessment: Gm32051, Mthfd2, Apol11b, Isg15, Rexo2, Atf5, Alad, Uba1, and Ank1 (Fig. 4A). Based on these results, we selected the top four genes: Gm32051, Mthfd2, Apol11b, and Isg15 (in descending order of expression fluctuations) (Fig. 4B, and C). In addition, we confirmed that the coverage graphs provided accurate read mappings (Fig. 4D). The four identified genes were quantified using the TB-Green qPCR assay to determine the reproducibility of the total RNA-seq data. The results showed that similar quantitative values for the total RNA-seq data were obtained with the expression values of the identified RNAs being more than 30–100-fold in AdEPO mice compared with control mice (Fig. 4E). Surprisingly, since these screened genes involved functions unrelated to EPO, these results suggest that even a single gene of the identified RNAs can be indirect proof of gene doping based on the concept of the ABP.
Figure 4. Single RNAs identified by bioinformatics screening as indirect proof. (A) Venn diagram showing the number of DEGs with filtering for high expression and high fold expression. (B) Scatter plot showing nine overlapping genes with filtering. (C) Relative gene expressions (Con. = 1.0) showing nine overlapping genes. (D) Genome browser indicating the read mapping results of the top four genes. (E) Conformation of the top four genes as indirect proof using the TB Green qPCR assay. *** \( p < 0.001. \)

3.4. Direct and indirect proof detected for up to approximately 30 days using a drop of whole-blood DNA/RNA

Fig. 5A shows an overview of the experiments performed in this section. The results showed that direct proof of viral genome DNA was significantly detected until 15 to 20 days after the injection (Fig. 5B-F). Furthermore, several mice qualitatively showed the target DNA fragments for the hEPO-1 and 2 primer-probe pairs (Pr-hEPO-1 and -2; Fig. 5B and C) until 30 days. Moreover, indirect proof of RNA expression levels identified from the total RNA-seq showed significant fluctuations, i.e., until only 5 days for Gm32051 as lncRNA and 20-30 days for other mRNAs (Fig. 5G-J). The longest time of significant changes was observed for Mthfd2 (Fig. 6H). These results indicate that direct proof of gene doping can be detected for a long time in association with changes in RNA expression as indirect proof.
Figure 5. Detection of direct and indirect proof for a long time period. To detect direct and indirect proof, small amounts of whole blood were harvested from the tail tips of mice (n = 12) until 30 days after the injection. From approximately 50 µL of whole blood, DNA/RNA were extracted, and the TaqMan or TB Green qPCR assays were performed to detect direct and indirect proof. (A) overview of the experiments in this section, (B) to (F) continuous detection of direct proof for 30 days by five primer-probe pairs using the TaqMan qPCR assay; “Pr” refers to a pri-
mer-probe, (G) to (J): detection of changes in gene expression as indirect proof using the TB Green qPCR assay. N.D.: not detected. *p < 0.05, **p < 0.01 and ***p < 0.001 vs. Pre value.

4. Discussion

The WADA has banned gene doping [3], and gene doping using the EPO gene, in particular, may be abused by athletes due to the recent rapid progression of gene therapy. However, detection methods for gene doping have not yet been established worldwide. Therefore, in this study, we first attempted to develop a mouse model of gene doping using an rAdV-hEPO vector. The gene and protein expression of hEPO in the liver were confirmed in AdEPO mice, indicating that the hEPO gene was successfully delivered to the liver. Moreover, secretion of hEPO protein as a hormone into blood were also confirmed in AdEPO mice. The gene expression of Gata1 (regulator of hematopoietic signaling) [31,32], Gypa, and Trfr (specific markers of RBCs) [33-35], were drastically increased in the spleens of AdEPO mice. Furthermore, the RBC count, Hgb level, and HCT were increased in AdEPO mice. We also supplementally clarified that these effects were not the result of injecting the rAdV-ZSG vector as hEPO-free rAdV. Therefore, these results indicate that the mouse model of gene doping made with the rAdV-hEPO vector was successfully established in this study. Moreover, the model may be useful to establish detection methods for gene doping.

Next, we aimed to develop detection methods for direct proofs of gene doping using the TaqMan-qPCR assay. The primers were designed to include the exon-exon junction of the hEPO gene, which can specifically detect transferred genes [6,24,25], because if the exogenous gene is transferred to organs, the intron DNA-sequences are not included. To detect robust proof of gene doping in mice, we also designed more primer-probes to target DNA sequences of CMVp, Hexon, and TKpA, which are specific to the rAdV-genome DNA sequence. All of the targeted DNA fragments could be detected, and the amplicons also had accurate DNA sequences as determined by Sanger sequencing. Therefore, these primer-probe pairs and qPCR conditions can be directly used to examine whole blood DNA from athletes. However, some caution is required. The gene cassettes of CMVp, Hexon, and Tkpa are applied on vectors for gene therapy [36]. Moreover, recently, the vaccine, named ChAdOx1 nCoV-19 (AZD1222), for SARS-CoV-2 may include the gene cassettes because the vaccine was made using replication-deficient simian adenoviral vector [37,38]. Therefore, individuals who have been treated with gene therapy or the vaccine may have false-positive test results, and determining whether subjects have received these treatments before conducting gene doping tests is necessary. In particular, if gene doping tests are performed at the 2020 Tokyo Olympics (held in July 2021), laboratories would need to confirm vaccination for SARS-CoV-2 in athletes because with the recent significant spread of COVID-19 by SARS-CoV-2 worldwide, athletes are more likely to travel to Japan after being vaccinated.

To investigate the period for which genome-specific DNA fragments of the rAdV-hEPO vector can be detected as proof of gene doping, the TaqMan qPCR assay was performed using DNA obtained from one drop of blood chronologically harvested from the mouse model. The statistical analyses showed that significant proof of gene doping could be detected until 20 days after injection of the vector. In addition, several mice showed positive results for viral fragments until 30 days, which can be used as the basis for a qualitative test. The metabolic rate of mice has been shown to be several times faster than that of humans [39]. Hence, in humans performing gene doping with the rAdV-hEPO vector, proof of doping may be detected in the blood for over a month, which is a longer time than in mice. Our previous study using droplet digital PCR reported that when the DNA extracted from one drop of whole blood from an rAdV gene doping mouse model was diluted and adjusted to a concentration of 10 ng/μL, significant positive proof of gene doping was detected until 5 days after the injection [5]. However, the present study showed that the use of undiluted DNA solution could provide significant proof of gene doping for a longer time than that reported previously. These results
suggest that undiluted DNA can be used to improve the sensitivity of detection. Therefore, we propose the use of undiluted DNA from whole blood samples in gene doping tests for humans.

As abovementioned, gene doping proof was detected using one drop of whole blood DNA. Currently, a small examination device for point-of-care testing (POCT) has frequently applied for human blood to examine several diseases at clinical sites [40]. As the technology of current POCT devices continues to improve, it may be possible to easily perform gene doping tests with PCR using one drop of blood collected from the fingertip after puncture with a small needle. Therefore, we have begun to develop a small examination device applied on the sports field for gene doping testing that could be less invasive. However, there are major barriers to this development. Hgb and IgG have been reported as factors that greatly inhibit PCR reactions [41]. In addition, a large amount of protein may also decrease the sensitivity of PCR. To develop an immediate POCT device for use on the sports field, it is necessary to resolve these issues, and we have repeated trial and error testing.

The ABP advocated by the WADA involves constant monitoring of selected biological variables that can indirectly reveal the effects of doping rather than attempting to detect the doping agent or method itself [20]. In other words, this concept is an attempt to detect traces of doping activity. The parameters evaluated as a part of ABP testing include general hematological parameters and steroid levels [20]. However, it is unclear whether RNA expression patterns in whole blood could be applied as parameters for the ABP. Here, we considered that total RNA-seq technology can be applied to identify RNAs as novel indirect proof based on the concept of the ABP because the peripheral blood transcriptome may dynamically reflect the biological mechanisms upon increased EPO secretion. Our results identified 1,128 DEGs in whole blood, and most of them were upregulated in AdEPO mice (Fig. 3B and C). The hematopoiesis-related terms such as GATA1 targets were significantly enriched in this profile by GSEA (Fig. 3D-F). These genes, such as erythrocyte surface proteins and erythroid transcription factors, are involved in erythropoiesis (Fig. 3G) [42-44]. For example, Tfrc encodes transferrin receptor protein 1 (sTfR), which is used for detecting rEPO hormone misuse by athletes and as a diagnostic test [45]. In addition, we predicted that RBC-specific genes were significantly enriched (Fig. 3H) and that other blood cell type-specific genes also were contained in DEGs (Fig. 3I). These results suggest that the expression changes, especially erythrocyte-specific genes, could reflect the hematopoietic effect of rAdV-hEPO and may serve as indirect proof based on the ABP. The enrichment analysis also showed that the erythropoiesis-related terms, such as mitochondria- and metal ion homeostasis-related terms, were significantly enriched in the upregulated genes (Supplementary Fig. S4A, C, E, and G.). For example, the deletion of Rb caused moderate anemia, induced inhibition of mitochondrial biogenesis, decreased OXPHOS pathway activity, and disturbed heme production and iron transport [46,47]. Thus, these results suggest that the blood transcriptome could explain and mirror the detailed hemopoietic mechanism induced by rAdV-hEPO injection.

We also performed TSEA to predict the tissue status using DEGs because the blood transcriptome and protein expression patterns can partly predict tissue status [48-50]. TSEA showed enriched hematopoiesis-associated tissues such as E14.5 liver, bone marrow, and spleen (Supplementary Fig. S5A). We noticed that the 42 predicted liver- and bone marrow-specific genes overlapped with each other, and these genes were associated with enriched hematopoiesis-related terms, such as porphyrin-containing compound biosynthetic process and erythrocyte development (Supplementary Fig. S5B-D). In particular, Gata1, known as the transcription factor for erythrocyte differentiation, was upregulated in blood, consistent with the findings for tissues (Supplementary Fig. S5E and F). These results suggested that the whole blood RNA profile may dynamically reflect the hematopoietic status in tissues injected with rAdV-hEPO.

Considering the results obtained to date, we believe that tracking significant changes in the overall profile of whole blood RNA expression can serve as a measure-
ment approach for ABPs since this idea is consistent with the concept of the ABP. Therefore, we recommend that total RNA-seq be employed in the generation of ABPs for gene doping in the future. However, to achieve this idea, it is essential to unify and establish a standardized workflow, such as RNA extraction, library preparation, NGS runs, and bioinformatics analyses, and a robust protocol with a low scope for artificial contamination during experimental operations.

Next, to identify single RNA as novel indirect proof based on the concept of the ABP, in the gene doping model, we screened potential single RNAs that showed high fold changes and high expression levels in the profile because the ideal genes should be measured with high-sensitivity qPCR. Interestingly, to our knowledge, the screened genes are not known to be directly associated with EPO function. In particular, we identified several RNAs whose expression drastically increased by 80- to 100-fold or more upon injection of rAdV-hEPO. The identified RNAs were both mRNAs and lncRNAs for genes such as Apol11b, Mthfd2, Gm32051, and Isg15 (Fig. 4). Apol11b (A330102K04Rik) is reported to share homology with the apolipoprotein L family of proteins in humans, suggesting its involvement in alterations in the lipid composition of erythrocyte membranes during terminal differentiation [51]. Mthfd2 is known as an enzyme involved in mitochondrial folate one-carbon metabolism and is also recognized as a NAD-dependent mitochondrial methylenetetrahydrofolate dehydrogenase-cyclohydrolase (NMDMC). Mthfd2 is reported to play a role in embryonic development and pluripotent stem cells in mice [52,53]. In addition, Mthfd2 is associated with a poor prognosis in hematological and solid cancers [54]. Isg15 is involved in modulating the development of erythroid differentiation [55]. To our knowledge, the function of Gm32051 remains unknown. Interestingly, the dramatically increased expression of these genes that have unknown function related to EPO may be new parameters for the ABP.

We also conducted long-term measurements of the identified RNAs as indirect proof, and the results showed that the expression levels of these RNAs significantly fluctuated for 15 to 30 days after injection of rAdV-hEPO. Therefore, these RNAs may be sensitive to injection of the rAdV-hEPO vector and could be used as novel parameters for the ABP. However, it is unclear whether the same results would be obtained in humans. Therefore, future studies should aim to conduct similar experiments with human participants. In addition, since it is also unclear why these RNAs sensitively responded to the rAdV-hEPO injection, an additional line of research should aim to elucidate the molecular level biological mechanisms in tissues and blood after injection of rAdV-hEPO.

Currently, automation of inspections is being actively pursued in the medical field. Therefore, human error is reduced, making testing more accurate. On the other hand, in this study, even though the control and AdEPO mice were completely separated during the experimental process, fragments of the Hexon region were detected in the blood of the control mice. Therefore, artificial cross-contamination could sway the results and must be avoided when conducting tests for humans. To avoid the abovementioned cross-contamination, we have tried to apply an automation system such as Maholo, which is an experimental robot similar to LabDroids [56], for gene doping testing. An automation system can perform its programmed contents with high precision in a clean environment free from artificial contamination. Therefore, an automatic system is expected to be effective in preventing deception on the part of the inspector, making the construct a more rigorous inspection system.

5. Conclusions

In summary, our study demonstrated that a gene doping mouse model was established using an rAdV-hEPO vector, and we showed that single parameters of the gene doping model including the concept of the ABP could be detected as direct and indirect proof of gene doping using one drop of whole blood DNA and RNA over a long term period. Moreover, we showed that the overall expression profiles of RNAs significantly changed in whole blood in the gene doping model and could become new parameters for
the ABP. These achievements provide new insights for the establishment of gene doping testing in human athletes. Therefore, the field of gene doping testing is entering the dawn of a new era.

**Supplementary Materials:** The following are available online at www.mdpi.com/xxx/s1, Figure S1: Blot raw data for the detection of total EPO in the liver, Figure S2: blot raw data for GAPDH expression in the liver, Figure S3: no hematopoietic effect of the rAdV unit alone in mice, Figure S4: GSEA predicting the detailed mechanisms of hematopoiesis induced by rAdV-hEPO in whole blood, Figure S5: TSEA predicting hematopoiesis in tissue status as reflected by gene expression profile, Table S1: primer and probe sequences.

**Author Contributions:** Conceptualization, methodology, validation, formal analysis, investigation, data curation, writing—original draft preparation, writing—review and editing, visualization, project administration, T.S., T.N. and S-i.F.; formal analysis, investigation, data curation, Y.M., G.I., K.A., K.Y., S.O. and S.T.; writing—review and editing, L.M. and H.U.; investigation, data curation, N.I. and K.T (Kenshiou Tamai); writing—review and editing, Y.K. (Yasuharu Kanki), Y.Y., K.W. and T.T.; writing—review and editing, supervision, Y.K. (Yasushi Kawakami) and K.T. (Kazuhiro Takekoshi). All authors have read and agreed to the published version of the manuscript.

**Funding:** This work was supported by grant of JAPAN Anti-Doping Agency in 2020.

**Institutional Review Board Statement:** The study was conducted in accordance with the guidelines (corporation regulation; No. 50, 21 July 2005) of the animal experiments by the Animal Ethics Committee of the University of Tsukuba. The approval number was 20-361 (1 May 2020).

**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** The total RNA-seq data have been deposited in the “Gene Expression Omnibus (https://www.ncbi.nlm.nih.gov/geo)” under accession number--------.

**Conflicts of Interest:** The authors declare no conflict of interest.
References

16. Sharma, R. Analytical study of doping cases of banned substances during Olympics games from 1968 to 2012. *Int J*


