Fluorometric evaluation of CYP3A4 expression using improved transgenic HepaRG cells carrying a dual-colour reporter for CYP3A4 and CYP3A7

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Primary human hepatocytes are necessary to evaluate cytotoxicity, drug metabolism, and drug–drug interactions for candidate compounds in early-phase drug discovery and development. However, these analyses are often hampered by limited resources and functional or genetic variation among lots. HepaRG human hepatocellular carcinoma cells can differentiate into mature hepatocyte-like cells (HepLCs) that possess similar metabolic activity to human hepatocytes. We previously established transgenic HepaRG cells carrying a dual reporter that expresses red fluorescent protein (RFP) under the transcriptional regulation of CYP3A7 in the hepatoblast-like cell state and enhanced green fluorescent protein (EGFP) under the transcriptional regulation of CYP3A4 following HepLC differentiation. In this study, we successfully isolated a subclone of transgenic CYP3A4G/7R HepaRG cells with an improved HepLC differentiation potency. Midazolam metabolism by CYP3A4 in these HepLCs was comparable to that in wild-type HepLCs. The EGFP fluorescence intensity was greatly induced by rifampicin (RIF) treatment. There was a strong correlation between fluorometric and metabolic analyses. The fold change in EGFP-positive cells was comparable to those in the CYP3A4 mRNA level and luminescence of proluciferin metabolites. RIF treatment and cell proliferation increased the RFP-positive cell number. Thus, CYP3A4G/7R HepLCs provide a real-time, multiwell-based system to co-evaluate CYP3A4 induction and hepatic regeneration.

Xenobiotic metabolism in vivo is mostly catalysed by cytochrome P450 isoenzymes (CYPs), which are mainly expressed in the liver and intestine1,2. CYP3A4 is involved in the oxidation of approximately 50–60% of drugs metabolized by CYPs3. Thus, the level of CYP3A4 enzymatic activity directly regulates the levels of metabolic reactions, leading to changes in the blood concentration of the compound itself and/or concurrent drugs. CYP3A4 is positively and negatively regulated in vivo through induction of its transcription and suppression of its enzymatic activity, respectively. These phenomena, termed CYP induction and inhibition, respectively, have a great impact on drug–drug interactions4. CYP3A4 expression is mainly induced through the heterodimer of nuclear receptor pregnane X receptor (PXR) and retinoid X receptor (RXR), which binds to the xenobiotic-responsive enhancer module located −7.8 kb upstream of the CYP3A4 transcription initiation site and proximal response elements5–7. PXR is activated by various compounds such as dexamethasone, rifampicin (RIF), and pregnenolone-16α-carbonitrile (PCN). Because of species specificity of metabolic enzymes and nuclear receptors caused by genetic differences, animal experiments cannot accurately evaluate drug–drug interactions in humans5,8. For example, mouse PXR is activated by PCN, but not by RIF. On the other hand, human PXR is poorly activated by PCN, but very effectively activated by RIF5,8. Moreover, CYP3A4 and PXR are...
mainly expressed in the liver and small intestine, but are not expressed in premature hepatic cells. Therefore, vast numbers of human adult hepatocytes are required in early-phase drug discovery and development.

Primary, cryopreserved, and long-term cultured human hepatocytes are used to predict reactions of compounds in humans. However, human liver cell resources are limited and their quality is variable. Differences in genetic background and environment among individuals also affect the accuracy and reproducibility of in vitro assays. In addition, human hepatocytes are only viable for a short period. Therefore, scalable adult-type hepatocyte-like cells (HepLCs) are globally desired.

Hepatic carcinoma and immortalized hepatocytes are homogeneous and proliferative; however, most of these cells poorly express CYP3A4. To overcome this issue, the human hepatocellular carcinoma cell line HepaRG has been studied. HepaRG cells are bipotent hepatoblast-like cells (HB-LCs) under proliferative conditions, dominantly express CYP3A7, a foetal liver-specific CYP3A isoform, and can differentiate into HepLCs and cholangiocyte-like cells. Metabolic activity in HepaRG-derived HepLCs is similar to that in human adult hepatocytes, in which the major CYP3A isoform is CYP3A4. The developmental shift of CYP3A isoforms mimics human perinatal development in vivo. Thus, CYP3A7 is a HB-LC marker, while CYP3A4 is an ideal marker of HepLCs.

Previously, we developed a procedure to restore the differentiation potency of a subclone of 4G/7R HepaRG cells based on the protocol used when the HepaRG cell line was originally established. In this protocol, we optimized the HepaRG differentiation protocol for the isolated subclone. 4G/7R HepaRG cells possessed a greatly improved HepaRC differentiation potency, and CYP3A4 enzymatic activity in HepaRCs derived these cells was comparable to that in WT HepaRC-derived HepLCs. The EGFP fluorescence intensity was obviously enhanced by CyP induction with Rif in 4G/7R HepaRCs, where the fold change in the EGFP fluorescence intensity strongly correlated with the CYP3A4 enzymatic activity. Thus, EGFP-positive 4G/7R cells are functional HepaRCs applicable for general purposes.

**Results**

**Restoration of the HepaRC differentiation potency in 4G/7R HepaRG cells.** WT HepaRG cells efficiently differentiated into HepaRCs upon culture in Medium 710 growth medium for 2 weeks and subsequent culture in Medium 720 differentiation medium for 2 weeks (Fig. 1A and 1B). By contrast, the original C3 clone of 4G/7R HepaRG cells was highly suscepLible to a high concentration of DMSO and mainly differentiated into cholangiocyte-like cells (Fig. 1B). When cells were seeded in Medium 710 at a density of about 5-fold higher than usual (2–3 × 10⁵ cells/cm²), all cells uniformly expressed RFP (Fig. 1C). To select cells resistant to DMSO, the DMSO concentration was gradually increased from 0.2% to 0.4% in Medium 710 during the 2-week growing culture period. Under these conditions, cells could continuously grow and express RFP, while some cells naturally differentiated into EGFP-positive HepLCS (Fig. 1D). During the course of subsequent differentiation culturing, we used Medium 710 containing 1.5% DMSO. After 2 weeks, RFP-positive cells disappeared and EGFP-positive cells became dominant (Fig. 1E). Differentiation of EGFP-positive cells was initiated by the disaggregation, and these cells became positive for RFP. After two rounds of growth culture in Medium 710 for 2 weeks followed by two rounds of stepwise differentiation for 4 weeks, we could finally isolate a subpopulation of 4G/7R HepaRG cells with HepaRC differentiation potency, as seen in WT HepaRG cells, based on cell morphology. In subsequent experiments, we gradually differentiated cells using a mixture of Medium 710 and Medium 720, as described in the Methods section.

Next, we compared the function of 4G/7R HepaRCs derived from the newly isolated subpopulation with that of WT HepaRG-derived HepLCS. Cryopreserved differentiated WT HepaRG cells are commercially available (Biopredic International). We cryopreserved differentiated 4G/7R HepaRG cells using a similar procedure. Defrosted differentiated cells were directly applied to all the assays presented here.
CYP3A4 Induction Tests

Fluoromicroscopic image analyses. We investigated whether the fold change in the total EGFP fluorescence intensity reflects that in the CYP3A4 transcription induction rate. Differentiated 4G/7R and WT HepaRG cells were defrosted and cultured in Medium 640 to eliminate DMSO from the cell culture environment because DMSO itself induces CYPs. Thus, the EGFP fluorescence intensity was greatly reduced in Medium 640, whereas cells were positive for red fluorescence and negative for green fluorescence, which represent the expression levels of CYP3A7 (CYP3A7-RFP) and CYP3A4 (CYP3A4-EGFP), respectively. 4G/7R cells were seeded at a high density and gradually differentiated in Medium 710 containing 0.2–0.4% DMSO for 2 weeks. The first round of differentiation was completed by additional culture for 2 weeks in Medium 710 containing 1.5% DMSO. Many EGFP-positive cells appeared, but they had not fully differentiated into HepLCs based on their morphology. The stepwise differentiation protocol was repeated three times. Scale bar, 100 μm.
This indicates two possibilities: RIF activates CYP3A7 transcription or the cytotoxic effect of RIF induces regeneration of HB-LCs (Fig. 2B). The CYP3A4 induction rate estimated based on the EGFP fluorescence intensity was 144.2 ± 49.4-fold higher in RIF-treated cells than in 0.1% DMSO-treated control cells (Fig. 2C, n = 3, p < 0.001).

Quantitative reverse transcription-polymerase chain reaction (RT-qPCR) analyses. The endogenous CYP3A4 mRNA level was evaluated by RT-qPCR. The CYP3A4 mRNA level was 8.7 ± 3.7-fold higher in RIF-treated cells than in 0.1% DMSO-treated control cells (Fig. 2D, n = 6, triplicate measurements, p < 0.001). In addition, the relative EGFP mRNA level was at least 3.9 ± 1.2-fold higher in RIF-treated cells than in 0.1% DMSO-treated control cells, when the EGFP mRNA was almost undetectable in 0.1% DMSO-treated control cells (Fig. 2D, n = 6, triplicate measurements, p < 0.001). These results demonstrated that fluorometric analyses of 4G/7R HepLCs are sensitive, leading to overestimation of the CYP3A4 induction rate by more than 10-fold compared with RT-qPCR-based estimations.

Flow cytometry (FCM) analyses. Approximately 2 × 10^4 living cells were quantitatively analysed using FCM (Fig. 3A). The frequency of EGFP-positive cells was 17.5 ± 3.6% in the 10 μM RIF-treated population and 1.5 ± 0.2% in the 0.1% DMSO-treated control population (Fig. 3A, n = 3). RIF treatment increased the CYP3A4 induction rate by 11.7 ± 2.4-fold based on EGFP-positive cell numbers (Fig. 3B, n = 3, p < 0.001). These results closely matched the findings regarding CYP3A4 mRNA expression levels, with a difference of only 1.3-fold.
Thus, EGFP-positive cell numbers may reflect CYP3A4 expression levels and can be used to evaluate CYP3A4 induction.

Liquid chromatography-tandem mass spectrometry (LC-MS/MS) analyses. CYP3A4 metabolic activity was compared between WT and 4G/7R differentiated cells by LC-MS/MS. A total of $0.5 \times 10^6$ differentiated cells were defrosted, resuspended in MEM containing 50 $\mu$M midazolam, and cultured for 1 hr. Then, the amount of the CYP3A4 metabolite 1′-hydroxymidazolam secreted by cells into the culture medium was determined. The level of 1′-hydroxymidazolam was 4.9 ± 1.0 nmol/hr/million cells for differentiated WT cells and 5.1 ± 0.8 nmol/hr/million cells for differentiated 4G/7R cells (Fig. 4A). These values were not significantly different, suggesting that the newly isolated 4G/7R subclone can differentiate into functional HepLCs at a high frequency similar to WT HepaRG cells. In the differentiated original C3 cells, the level of 1′-hydroxymidazolam was almost undetectable (0.04 ± 0.00 nmol/hr/million cells) and was significantly lower than in WT and 4G/7R cells (Fig. 4A, n = 4, p < 0.001). Thus, CYP3A4-mediated metabolic function is considerably restored in the newly isolated 4G/7R subclone.

Next, the CYP3A4 induction rate was evaluated based on the fold change in the level of 1′-hydroxymidazolam. Cells were treated with 10 $\mu$M RIF or 0.1% DMSO in Medium 640 for 48 hr. Midazolam was added to the medium for the final hour of cell culture. Then, the supernatant was analysed by LC-MS/MS. The level of CYP3A4 metabolites was 6.7 ± 2.1-fold higher in differentiated WT HepaRG cells and 13.3 ± 7.0-fold higher in differentiated 4G/7R cells than in 0.1% DMSO-treated control cells (Fig. 4B, n = 3, p < 0.001 and p < 0.01, respectively). These two values did not significantly differ. In addition, there was a strong positive correlation between the CYP3A4 induction rates estimated by the EGFP fluorescence intensity and the amount of 1′-hydroxymidazolam (Fig. 4C, r = 0.95) when the two estimation methods were sequentially applied to the seven sets of independently differentiated cell samples. However, the CYP3A4 induction rate estimated based on the EGFP fluorescence intensity was 10-fold higher than the rate estimated by the amount of metabolites. These results indicate that EGFP images captured by fluorescence microscopy provide a simple, quick, and sensitive system to evaluate CYP3A4 induction.

Luminometric CYP3A4 induction tests using a proluciferin substrate. Relative CYP3A4 metabolic activity was measured based on the fold change in luminescence using proluciferin IPA (Promega) as a CYP3A4-specific substrate. Luminescence was 10.6 ± 1.0-fold higher in 10 $\mu$M RIF-treated 4G/7R cells than in 0.1% DMSO-treated control cells (Fig. 4D, n = 6, p < 0.001). These results closely matched the findings regarding EGFP-positive cell numbers estimated by FCM, with a difference of only 0.9-fold.

Discussion
In this study, we restored the HepLC differentiation potency of previously established transgenic 4G/7R HepaRG cells. We also developed a differentiation method optimized for this subclone. The transgenic 4G/7R subclone possesses the following properties: (1) undifferentiated cells are HB-LCs positive for RFP under the control of
enhancer and promoter regions of CYP3A7; (2) differentiated cells become negative for RFP; (3) HepLCs become positive for EGFP under the control of enhancer and promoter regions of CYP3A4; (4) CYP3A4 enzymatic activity in differentiated 4G/7R cells is equivalent to that in differentiated WT HepaRG cells; (5) the fold change in the EGFP-positive cell number is similar to those in the levels of CYP3A4 metabolites and CYP3A4 mRNA in RIF-treated cells, and (6) the fold change in the EGFP fluorescence intensity is 10-fold higher than those calculated in other assays.

CYP3A4 induction often causes drug–drug interactions in clinical practice. Therefore, it is necessary to predict the CYP3A4 induction potency for numerous candidate compounds in the research phase of drug discovery and development. HepaRG cells exhibit a liver-specific function and can provide many differentiated HepLCs, which can be cryopreserved for long periods. In this study, we isolated a 4G/7R cell subclone that can provide many CYP3A4-expressing HepLCs, in which CYP3A4 transcriptional and enzymatic activities can be easily predicted based on EGFP fluorescence. CYP3A4 activities can be evaluated using LC-MS/MS, FCM, and RT-qPCR, which are time-consuming. However, these assays are not applicable to multiwell-based screening.

In early-phase drug discovery and development, high content screening (HCS)-based analyses are required to investigate the properties of numerous candidate compounds simultaneously. 4G/7R HepLCs may be suitable for multiwell-based HCS analyses based on the EGFP fluorescence intensity. The EGFP fluorescence intensity-based assay overestimated the CYP3A4 induction rate by 10-fold. This is advantageous to prohibit false-negative judgments in the initial screening. In addition, the number of HepLCs applied to each well can be reduced.

In mice and rats, three small molecules, Y-27632, A-83-01, and CHIR99021, were recently identified that induce hepatic regeneration in vitro and can convert mature hepatocytes into bipotent HB-LCs, termed chemically induced liver progenitors. It is important to identify such small molecules to induce hepatic regeneration in humans after hepatic injury. In this study, reactivation of RFP was observed when EGFP-positive HepLCs dedifferentiated, and then RFP-positive cells became EGFP-positive HepLCs again. Thus, RFP fluorescence can be used as a marker of hepatic regeneration caused by hepatotoxicity, together with erasure of EGFP. Therefore, we believe that 4G/7R HepLCs will also be useful for an HCS-based assay to eliminate hepatotoxic drugs and to identify hepatic regenerative medicines.

Some hepatocyte-specific genes related to CYP and transporter activities are poorly expressed in differentiated WT HepaRG cells. This is one reason why the general use of HepaRG cells is limited. Further gene manipulation will resolve this problem by enhancing expression of a set of hepatocyte-specific genes in HepaRG cells. Our culturing procedure will be applicable to generate ideal HepLCs from HepaRG cells as scalable and uniform model cells of human primary hepatocytes.

**Methods**

**Cell culture.** Undifferentiated 4G/7R HepaRG cells were seeded at a density of 1.5 × 10^5 cells/cm² and cultured in Medium 710 for 2 weeks at 37°C in 5% CO₂. The medium was changed once every 3–4 days. For differentiation, cells were cultured for 6 days in a mixture of Medium 710 and Medium 720 at a ratio of 3:1 and then further cultured for 2 days in a mixture of these media at a ratio of 1:1. Then, differentiation culture was performed for 12 days in Medium 720. Differentiated cells were stored in liquid nitrogen. The differentiated cell population was subjected to each of the analyses to investigate correlations among the results. All media used for HepaRG cell culture were purchased from Biopredic International.
5 Goodwin, B., Hodgson, E. & Liddle, C. The orphan human pregnane X receptor mediates the transcriptional activation of CYP3A4 with 342.1(Q1)/168.0(Q3) for 1 exit potential. Detection with the mass spectrometer was performed using multiple reaction monitoring mode with the source gas 2, 101 V declustering potential, 10 V entrance potential, 51 V collision energy, and 10 V collision cell plate, and cultured in Medium 670 for 96 hr at 37 °C in 5% CO2. During the final 48 hr of culture, cells were cultured in Medium 640 containing 10 μM RIF or 0.1% DMSO. For LC-MS/MS analyses, cells were incubated in MEM containing 50 μM midazolam at 37 °C during the final 1 hour of culture. RIF and midazolam were dissolved in DMSO and stored at −20 °C.

FCM analyses. Cells treated with 10 μM RIF or 0.1% DMSO for 48 hr were suspended in Medium 710, and the frequencies of EGFP- and/or RFP-positive cells were analysed using a Gallios Flow cytometer (Beckman Coulter). EGFP and RFP were excited with a 488-nm blue laser and detected using 525-nm and 575-nm band-pass filters, respectively.

LC-MS/MS analyses. Differentiated cells were thawed in Medium 670 and incubated for 1 hr in MEM containing 50 μM midazolam at 37 °C in 5% CO2. The culture supernatant was diluted with methanol at a ratio of 1:3, and cell debris was removed from the sample by centrifugation at 15,000 rpm for 15 min at 4 °C. The supernatant was diluted with water containing 3 μM α-hydroxymidazolam-D4, an internal standard. The levels of 1′-α-hydroxymidazolam and α-hydroxymidazolam-D4 were quantified using LC-MS/MS. The liquid chromatography experiments were conducted with a prominance UFLC system (SHIMADZU) coupled with QTRAP5500 (SCIEX). Chromatographic separation was achieved in a TSK gel ODS-100V column (50 mm × 0.2 mm, 3 μm, Tosoh) at 40 °C with an injection volume of 10 μl. The mobile phase consisted of solvent A (0.1% formic acid prepared in water) and solvent B (0.1% formic acid prepared in acetonitrile). The flow rate was 0.2 ml/min. The isocratic specification conditions were used with a composition of A:B = 80:20 (v/v). Mass spectrometry was performed using electrospray ionization in positive ion mode under the following operating conditions: 40 psi curtain gas, 9 collision gas, dimension-less, 5500 V ion spray voltage, 650 °C, 30 psi ion source gas 1, 40 psi ion source gas 2, 101 V declustering potential, 10 V entrance potential, 51 V collision energy, and 10 V collision cell exit potential. Detection with the mass spectrometer was performed using multiple reaction monitoring mode with 342.1(Q1)/168.0(Q3) for 1′-α-hydroxymidazolam and 346.1(Q1)/169.2(Q3) for [2H4]-hydroxymidazolam. M/Z 342.1 and 346.1 were molecular ions ([M + H]+) of 1′-α-hydroxymidazolam and [2H4]-hydroxymidazolam, respectively. Data were acquired using Analyst 1.5 software (SCIEX).

Microscopic image analyses. Fluorescence microscopic images were captured with a constant exposure time (Nikon). The mean and standard deviation of the total area fluorescence were calculated from the images using Image J software (an open-source program for image analysis provided by the National Institutes of Health).

RNA extraction and RT-qPCR analyses. Total RNA was extracted from 4G/7R cells using the RNeasy Mini Kit (Qiagen) and used to synthesize cDNA using the Superscript III First-Strand Synthesis Kit (Invitrogen). cDNA derived from 10 ng total RNA was amplified in a 25 μl reaction using the Power SYBR Green PCR Master Mix Kit (Applied Biosystems) and a LightCycler 480 system (Roche Applied Science). The primers used are described in Table 1.

Luminometric analyses using Luciferin IPA. CYP3A4 enzymatic activities were measured by a luciferase-mediated assay using the P450-Glo™ CYP3A4 Assay Luciferin-IPA kit (Promega) according to the manufacturer’s protocol. Luminescence was detected with the plate reader Infinit F500 (Tecan).

Statistics. Data are presented as means ± standard deviation. The Mann-Whitney U-test statistically compared the treated and control samples. Significance was determined using two-tailed equal variance Z values. p < 0.01 was considered significant.

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Author Contributions
S.T., M.T., and T.S. designed and performed the experiments. S.T. and M.T. performed the computational analysis. T.U. and M.T. interpreted the data and wrote the paper. M.T. initiated and directed the project.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

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