Autofluorescence in freshly isolated adult human liver sinusoidal cells

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Autofluorescent granules of various sizes were observed in primary human liver endothelial cells (LSECs) upon laser irradiation using a wide range of wavelengths. Autofluorescence was detected in LAMP-1 positive vesicles, suggesting lysosomal location. Confocal imaging of freshly prepared cultures and imaging flow cytometry of non-cultured cells revealed fluorescence in all channels used. Treatment with a lipofuscin autofluorescence quencher reduced autofluorescence, most efficiently in the near UV-area. These results, combined with the knowledge of the very active blood clearance function of LSECs support the notion that lysosomally located autofluorescent material reflected accumulation of lipofuscin in the intact liver. These results illustrate the importance of careful selection of exogenous fluorophores, especially when labelling of live cells where the quencher is not compatible.

Key words: Autofluorescence; endogenous fluorophores; liver; endothelial cells; lipofuscin.

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Introduction

Autofluorescence, the natural emission of light by biological structures, may possibly hamper fluorescence-based techniques if not properly addressed and corrected for. Several endogenous fluorophores are known to cause auto fluorescence in many tissues, including liver. Auto fluorescence in normal human liver tissue is predominantly caused by NAD(P)H, while vitamin A (vit A) is known to contribute to the auto fluorescence observed in hepatic stellate (Ito) cells. The fluorescent pigment lipofuscin accumulates in the lysosomes of many cell types with age and/or in patients with severe malnutrition and cancer cachexia and due to its broad excitation and emission spectra its presence complicates the use of techniques employing exogenous fluorophores. While most endogenous fluorophores have distinct spectral emission profiles, the spectra of lipofuscin overlap those of all commonly used exogenous fluorophores. This makes it difficult or even impossible to distinguish between specific labelling and auto fluorescence caused by lipofuscin.

Liver sinusoidal endothelial cells (LSEC) rapidly change their morphology and function in culture and cell lines which retain the in vivo characteristics of these sinusoid lining cells are not yet available. Studies on the physiological function of these cells thus depend on isolation of primary cells from tissue samples. Several methods depending on fluorescence are often used to isolate, characterize, and investigate primary liver cells from human and animals. While auto fluorescence in hepatocytes and stellate cells is well known, the presence of endogenous fluorophores in human LSEC is little known. In this study we aimed to characterize the observed auto fluorescence in human LSEC and determine whether pretreatment with a lipofuscin auto fluorescence quencher would allow for subsequent use of exogenous fluorophores.

Materials and Methods

Human liver tissue and isolation of human liver sinusoidal endothelial cells

Liver tissue was obtained from patients undergoing hepatic resection at the Department of Clinical Science, Intervention and Technology (CLINTEC), Karolinska University Hospital, Stockholm, Sweden or the University Hospital of North Norway (UNN), Tromso, Norway. Patients included in the study gave informed consent and ethical approval of the study was granted by the Regional Committee for Medical and Health Research Ethics (REC ID 7065). Perfusion of human liver tissue resections has previously been described in detail. Only minor modifications were implemented, as described below. After cannulation, the tissue was washed using cold Ca²⁺- and Mg²⁺-free Hanks Balanced Salt Solution (HBSS) followed by perfusion with warm (37°C) HBSS with 50 μg/mL ascorbic acid and 0.5 mM EGTA. The tissue was then perfused with warm HBSS without additives to remove EGTA before 25-30 mg/L liberase TM (cat.no: 05401127001; Roche, Basel, Switzerland) in 200 mL CBC buffer (0.4% w/v NaCl, 0.025% w/v KCl, 2.3% w/v Heps, 0.26% w/v NaOH, 0.07% w/v CaCl₂) was recirculated until optimal tissue digestion. The cell suspension was filtered through 100 μm sterile mesh and hepatocytes pelleted by centrifugation at 50 g for 3 min at 4°C. Non-parenchymal cells (NPCs), pelleted by centrifugation at 300 g for 5 min at 4°C, were fractionated using a 25% / 50% Percoll (cat.no: P1644; Merck, Darmstadt, Germany) gradient (1350 g, 25 min, 4°C). The cells located at the interface between the two centrifugations of Percoll were subjected to positive selection using CD31+ magnetic microbeads (cat.no: 130-091-933; Miltenyi Biotec, Auburn, CA, USA). The images shown in the manuscript originates from 3 different patient samples, but auto fluorescence was observed in all samples originating from adult humans undergoing hepatic resections due to various cancer diagnoses.

Fluorescence and confocal microscopy

CD31+ liver cells, enriched in LSEC, were resuspended in AIM-V medium (cat.no: 12055091; Gibco, Amarillo, TX, USA) and plated in 8 well ibidi µ-slides coated with quick coating solution (cat. no: cAP-01, Angio-Proteomic; Boston, MA, USA) at a density of 150,000-250,000 cells/cm². The cells were incubated at 37°C in 5% CO₂ and 3% O₂. Cells were fixed for 30 min at room temperature (RT) using 4% buffered formaldehyde (FA) after 12-48 h in culture. Confocal images were captured using a Zeiss LSM800 system equipped with a 40X 1.2NA water immersion lens before and after treatment with TrueBlack™ Lipofuscin Auto fluorescence Quencher (cat.no: 23007; Biotium Inc., Fremont, CA, USA). Live and fixed cells were also imaged in a Nikon fluorescence microscope equipped with a Zeiss AxioCam.

Imaging flow cytometry

CD31+ LSEC enriched liver cells in suspension were lightly fixed in 4% FA for 5 min. The cell suspension (20 million cells/mL) was transferred to 5 mL tubes with cell strainer caps (BD Biosciences) before analyzed on an AMNIS® ImageStream® MKII imaging flow cytometer (Luminex). Excitation lasers at 405 nm, 488 nm, 561 nm, and 642 nm were included to simultaneously capture fluorescent images in the emission range of 445-772 nm of in total 200 cells/patient.

Immunolabelling

Cultured CD31+ LSEC enriched liver cells were fixed as previously described, permeabilized with 0.2% Triton X-100 in PBS for 4 min and treated with TrueBlack™. Immune labelling was performed using goat anti-human mannose receptor (hMR) antibody (AF2534, R&D systems, 4 μg/mL), mouse anti-human Fc gamma receptor II (FcɣRII) (FUN2, Biolegend, 5 μg/mL), mouse anti-human LAMP-1 antibody (H4A3, Developmental studies hybridoma bank, 0.3 μg/mL). Primary antibodies were diluted in 1X PHEM buffer (60 mM PIPES, 25 mM HEPES, 10 mM EGTA, and 4 mM MgSO₄·7H₂O) containing 1% bovine serum albumin (BSA) and incubated for 1 h at RT. Secondary antibodies were Alexa Fluor (AF) 488 donkey anti-goat IgG and AF488 donkey anti-mouse IgG (Invitrogen, 4 μg/mL), mouse anti-human Fc gamma receptor II (FcɣRII) (FUN2, Biolegend, 5 μg/mL), and mouse anti-human LAMP-1 antibody (H4A3, Developmental studies hybridoma bank, 0.3 μg/mL). Primary antibodies were incubated in 1X PHEM buffer (60 mM PIPES, 25 mM HEPES, 10 mM EGTA, and 4 mM MgSO₄·7H₂O) containing 1% bovine serum albumin (BSA) and incubated for 1 h at RT. Secondary antibodies were Alexa Fluor (AF) 488 donkey anti-goat IgG and AF488 donkey anti-mouse IgG (Invitrogen, 4 μg/mL), diluted in 1X PHEM containing 1% BSA and incubated for 30 min at RT. The fluorescent DNA stain DAPI (10 mM), incubated for 15 min at RT, was used for visualization of nuclei.

Transmission electron microscopy

Cultured cells were fixed in 4% FA, 1% glutaraldehyde (GA) and prepared as described by Kuo. In brief, fixed cells were scraped and pelleted before washing in 1X PHEM buffer, post fixed in 1% aqueous Osmium Tetroxide for 1.5 h and dehydrated in a graded series of ethanol, with dry acetone as an intermediate step, before infiltration in an Epon substitute [AGAR 100 resin (R1043), DDSA (R1051), MNA (R1081) and DMP-30 (R1065)] and polymerized at 60°C overnight. Ultrathin sections were made on a Leica EM UC7 Ultramicrotome using a Diatome diamond knife, mounted on carbon-coated formvar films on copper grids and contrasted with 5% Uranyl acetate and Reynold’s lead citrate. Images were taken using a Hitachi HT7800 microscope equipped with an Emsis Xarosa digital camera.
Results

A large proportion of freshly isolated live and fixed CD31+ enriched LSEC displayed strongly fluorescent granules in the cytoplasm, appearing in all channels of both conventional (RGB) and confocal fluorescence microscopy (405 nm/blue, 488 nm/green, 561 nm/red, 633/far red) using normal laser intensity (Figure 1 B,C), although with less magnitude in the blue channel. The autofluorescent material was mostly located in the perinuclear area and appeared both as larger (up to 5 μm in diameter) and smaller (<1 μm in diameter) granules. Imaging flow cytometry confirmed that the autofluorescence was detected in all available channels (Figure 1D).

Figure 1. Autofluorescence in primary human LSECs. A) Conventional brightfield image of freshly isolated CD31+ LSECs in culture 17 h after plating; scale bar: 50 μm. B) Same area of cells imaged in a Nikon fluorescence microscope using brightfield (top), green filter (middle) and red filter (bottom); scale bar: 100 μm. C) A large proportion of non-treated CD31+ LSECs labelled only with DAPI shows autofluorescence in all four channels (blue, green, red, and far red) at normal laser intensity; scale bar: 10 μm; to watch a video of the autofluorescence through the entire spectrum of a Zeiss confocal microscope equipped with diode, Argon, DPSS and HeNe lasers, please visit Google drive at https://tinyurl.com/CD31autofluorescence. D) A large proportion of freshly isolated, non-treated, and lightly fixed CD31+ cells show autofluorescence in the 12-channel layout using AMNIS® imaging flow cytometry; the image is a representative view of 200 cells imaged.
Time-dependent treatment with TrueBlack™ strongly reduced the autofluorescence, most efficiently in the near UV-area (413-483 nm) (Figure 2). When applied for 2 min, TrueBlack™ efficiently quenched autofluorescent signals when applying moderate laser intensity (0.5% laser gain, <700 mV amplitude) to the samples. However, when increased laser intensity was applied, the autofluorescence reappeared.

Subsequent immunostaining for specific cell markers was not impaired by the TrueBlack™ treatment. Immune labelling showed that the autofluorescent granules were present in cells that stained positively for two human LSEC markers, FcɣRII (expressed in zone II/III) and the mannose receptor (Figure 3A).14-17 The perinuclear localization suggested that the granules could be associated with structures belonging to the lysosomal compartment and immunostaining revealed that the immunofluorescent material was located within LAMP-1 positive vesicles (Figure 3B).

Discussion

Autofluorescence in liver tissue is well known,1 but the morphological appearance and spectral characteristics of the well-established endogenous liver fluorophores NAD(P)H and vit A do not match the features of the autofluorescent granules observed in CD31+ LSECs. Vit A-induced autofluorescence in murine stellate cells is visible as large lipid droplets in the perinuclear area with increasing size in older animals,18,19 but in human cells it appears as smaller droplets scattered throughout larger areas of the cytoplasm.20 Furthermore, the spectrum of vit A-induced autofluorescence resides in the range of excitation 370-380 nm/emission 490-510 nm (blue region).6 In contrast, the autofluorescent granules in the CD31+ LSECs were visible throughout the entire emission range. Additionally, vit A-induced autofluorescence has high photolability and rapidly bleaches when exposed to light.1 The autofluorescent granules of the CD31+ LSECs did not show any signs of bleaching when excited with different light during fluorescence or confocal imaging.

LSECs have a high endocytic capacity and are known for their
scavenging ability, removing macromolecular waste products from extracellular matrix turnover from the blood circulation. This, along with the fact that lipofusins are remnants of tissue turnover products, suggests that LSECs accumulate this substance. In addition, a number of pathological conditions, including cancer, are associated with the accumulation of lysosomal pigments (ceroid-type lipofuscin) displaying properties similar to lipofuscin. The liver tissue samples included in this study originated from adult patients with metastatic colon cancer (part of the disease-free resection margin) and the possibility of increased lipofuscin accumulation in cells isolated from this type of tissue cannot be ruled out.

The pH indicator phenol red, present in many cell culture media, is fluorescent when excited at 440 nm. Accumulation of phenol red in lysosomes could occur during culturing of the cells. The cells examined in the present work have been kept in culture for only a short period of time (12-24 h) and the cells analysed by the AMNIS® imaging flow cytometry were not cultured, supporting the notion that the autofluorescent material represents lipofuscin.

Although the autofluorescence can be quenched, there are some technical issues arising from this phenomenon. Quenching of the autofluorescence works best when applying normal laser intensities and will reappear if the laser intensity increases. When work-

![Figure 3. Autofluorescence and exogenous fluorophore detection in immunolabelled LSEC-marker expressing cells. A) CD31+ LSECs were cultured for 24 h, fixed, treated with TrueBlack™ for 2 min and immunolabelled using (left) mouse anti-human FcγRII (green) and (right) goat anti-hMR (green) before being imaged using moderate laser intensity (0.5% laser gain, 700 mV amplitude); the intensity of the red laser was increased to 1% laser gain, 780 mV amplitude for the autofluorescence to reappear after TrueBlack-quenching (red); nuclei are stained with DAPI; scale bar: 20 μm. B) Lysosomal labelling with mouse anti-human LAMP-1 antibody (green) shows that the autofluorescent material (red) is located within LAMP-1 positive vesicles (arrow heads); cells were treated with TrueBlack™ for 1 min prior to immunolabelling to quench autofluorescence in the green channel while keeping some autofluorescence in the red channel; scale bar: 10 μm. C) TEM micrograph of a human LSEC displaying large perinuclear vacuoles (arrows) filled with material, displacing an indented (arrowheads) nucleus (N); scale bar: 2 μm.](image-url)
ing with low abundancy targets, this may introduce a conflict between visualizing the target and reappearance of the autofluorescence. Additionally, the lysosomal location of the autofluorescence may create technical challenges if cellular trafficking involving the lysosomal compartment is to be investigated.

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