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Clarification and confocal imaging of the nonhuman primate placental micro-anatomy

JA Sargent¹,², VHJ Roberts², JE Gaffney² & AE Frias¹,²

ABSTRACT

Geometry of the placental villous vasculature is a key determinant of maternal–fetal nutrient exchange for optimal fetal growth. Recent advances in tissue clarification techniques allow for deep high-resolution imaging with confocal microscopy; however, the methodology lacks a signal-to-noise ratio of sufficient magnitude to allow for quantitative analysis. Thus, we sought to develop a reproducible method to investigate the 3D vasculature of the nonhuman primate placenta for subsequent data analysis. Fresh placental tissue was dissected, formalin fixed, clarified using a modified Visikol® protocol and immunolabeled for CD31 (fetal endothelium) and cytokeratin-7 (villous trophoblast) for confocal imaging of the microanatomy. We present a detailed clarification and staining protocol augmented for imaging of nonhuman primate placental tissue. The image stacks generated by this refined staining method and our data acquisition parameters can be analyzed quantitatively to provide insights regarding the villous and vascular microanatomy of the placenta.

METHOD SUMMARY

We present a protocol for the preparation, immunolabeling and clarification of the nonhuman primate placenta refined for confocal microscopy and subsequent quantification of the microanatomic structures.

KEYWORDS

Confocal microscopy • Immunohistochemistry • Nonhuman primate • Placenta • Tissue clarification

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Through the imaging of immunofluorescent-labeled and clarified tissue with a confocal microscope, new approaches are emerging that address the shortfalls of stereological and vascular casting methods. The use of confocal microscopy to capture precisely located optical sections allows for the 3D rendering of the images [18–20]; however, the depth of tissue analysis has been limited due to refractive index mismatching, subsequent light-scattering, and auto-fluorescence [21]. The clarification of biologic specimens through refractory index equilibration has been studied since the beginning of the 20th Century, however earlier methods were damaging to tissue and incompatible with fluorescently labeled specimens [22]. To take advantage of the increasingly sophisticated confocal methodologies available, newer tissue clarification methods have been developed that are either based on hydrogel embedding, hyper-hydration, simple immersion or solvent-based techniques; each of these having their respective deleterious effects on tissue structure, or compatibility issues with immunofluorescently labeled tissue [22]. Visikol® HISTO™ is a proprietary reversible clearing technique that uses an ethanol-based solvent, which causes minimal tissue alteration (shrinkage/expansion of <5%) and is compatible with immunofluorescent staining. Recently, Visikol HISTO has been utilized to generate 3D renderings of immunofluorescent-labeled and clarified human placental tissue [23,24], and this has generated interest in customizing its use for novel analyses. Unfortunately, protocols refined for human application did not yield stacks of sufficient quality so as to allow for subsequent reliable quantitative analysis when applied to the nonhuman primate placenta. Specifically, chemical tissue fixation with Zn Formalin is required to ensure microanatomic structure stabilization during the long incubation times required for deep tissue.
imaging. The subsequent molecular cross-linking masks tissue antigens that are of scientific interest, and antigen-specific antigen retrieval steps must be optimized in order to allow for successful immunofluorescent labeling. Additionally, when exposed to laser excitation, many biologic tissues contain molecules (e.g., elastin, collagen, lipofuscin and heme byproducts) that fluoresce within the visible wavelengths (350–630 nm); if not reduced, this auto-fluorescence creates optical noise and interferes with quantitative analyses. Photo-bleaching of natural fluorophores can be accomplished in a wavelength-specific manner through photo-induced free-radical generation and chemical destruction using light-emitting diodes (LEDs) without disrupting the tissue integrity [25]. This has not been achieved in placental confocal imaging studies to date.

Numerous computer programs have been created to process confocal images and allow for image optimization, 3D rendering, and subsequent analysis. ImageJ is a versatile, Java-based, image-processing, free software developed by the NIH, which has a flexible development model allowing for simple user-generated plug-ins to increase the programs functionality [18,26]. The 3D image-based quantification of vascular networks has been approached in various fields of medicine [27], and ImageJ has a multitude of plugins that have been validated for this use making it suitable for use in our study [28].

Placental and fetal development is similar between humans and nonhuman primates (NHPs), and NHPs have been used successfully as a translational model to human pregnancies. Numerous validated, well-characterized, highly controlled NHP pregnancy disease models exist providing a unique opportunity to assess underlying disease mechanisms and potential therapeutic strategies. Using confocal imaging methods, quantified analysis of the NHP placenta has not yet been accomplished; however, this could provide critical insights into both human and NHP pregnancies. Our objective was to develop and refine a protocol for the immunofluorescent labeling and clarification of the placenta in the NHP to allow for high-resolution 3D imaging of sufficient quality for advanced quantification methods (Figure 1).

### MATERIALS & METHODS

Figure 1 provides a schematic overview of the seven stages of our protocol as outlined in detail below.

#### Sample preparation

All animal procedures were conducted in accordance with the guidelines of the Institutional Animal Care and Use Committee (IACUC) of the Oregon National Primate Research Center (ONPRC). The ONPRC abides by the Animal Welfare Act and Regulations enforced by the United States Department of Agriculture. Adult female rhesus macaques underwent time-mated breeding to generate pregnancies from which placental tissue was obtained. All animals were pair housed and maintained on a control standard chow diet with additional access to daily enrichment foods and foraging devices. All placental tissues were obtained under general anesthesia at the time of cesarean section delivery at gestational day 140 (term being 168 days), immediately placed in phosphate-buffered saline (PBS), and the decidua and membranes dissected away from the villous tissue. 1 mm³ tissue sections were isolated and immersed in 10% zinc formalin for 4 h prior to transfer in 70% ethanol and storage at 4°C. For the development of this protocol we utilized over 300 samples from 20 NHP pregnancies. Shorter formalin exposure times were associated with incomplete fixation, while longer incubations resulted in proportional worsening of the background auto-fluorescence.

#### Removal of heme byproducts

To address background fluorescence caused by hemoglobin byproducts, fixed tissue was placed in PBS and gently rocked at 4°C (Boekel Rocker II 260350–2 Platform Rocker, Boekel Scientific, PA, USA) for 48 h. Tissue was dehydrated with 50, 75 and 100% methanol washes with gentle rocking at 4°C for 10 min per wash. Using a bleaching solution (30% H₂O₂, dimethyl sulfoxide [DMSO], methanol in a 1:1:4 ratio), tissue was incubated at 4°C for 48 h under gentle rocking. We found that the 48-h exposure times, with agitation, were required to remove any retained hemoglobin byproducts, which otherwise create excessive background noise and artifacts.

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unacceptably interfere with quantitative analysis. To reduce general auto-fluorescence, heat exposure was limited and all steps were performed under LED bulbs with wavelengths of 480 nm and 660 nm (45W Multi-Spectrum LED, iPower, CA, USA) at a distance of 6 cm.

Antigen retrieval
For immunofluorescent staining, antigen retrieval was essential. Tissue was rehydrated using 20% DMSO/methanol, 75 and 50% methanol, and PBS washes with gentle rocking at 4°C for 10 min per wash. Optimal antigen retrieval was obtained using Citrate Buffer (10 mM citric acid with 0.05% Tween at pH 6) exposed to 80°C at 6 psi (Cuisenart Electric Pressure Cooker – CPC600, Cuisenart, NJ, USA) for 40 min, and then placed at 21°C for 20 min. Antigen retrieval times of 10, 20, 40 and 60 min were tested and heat and pressure conditions were varied. While the CK7 immunofluorescent labeling was robust over the breadth of trials, the final exposure conditions were required to optimize the CD31 labeling while minimizing the worsening background auto-fluorescence caused by the heat and pressure conditions. Given the specific optimization required for our two antigens, it is likely that a variety of conditions will need to be tested in order to modify this protocol if alternative structural labels are explored by this method.

Immunofluorescent labeling
Blocking and permeabilization was performed using a solution of 2% Donkey Serum and 0.2% Triton X-100 in PBS with gentle rocking at 4°C for 24 h under LED exposure at 4°C. A primary antibody solution containing both a monoclonal mouse-anti-human CD31 (Thermofisher MA1–26196, endothelial cell marker) and a polyclonal rabbit-anti-human CK7 (Abcam, ab103687, trophoblast marker) was made at 1:10 concentrations in a solution of 2% donkey serum in PBS, and tissue was immersed and placed at 37°C for 72 h. Four 10 min washes of 2% Donkey Serum in PBS at 21°C were then performed before placing in a secondary antibody solution of donkey-anti-mouse pre-adsorbed polyclonal antibody (Abcam, ab150109, alexafluor 488) and a donkey-anti-rabbit antibody (Abcam, ab150075, alexafluor 647) at a concentration of 1:250 for both secondaries in 2% Donkey Serum and PBS for 12 h at 21°C with agitation at 150 rpm. Shorter incubation times and altered temperatures (4, 21 and 37°C were trialed) were associated with incomplete tissue penetration as assessed by subjective observation at the time of imaging where antibody-positive staining was visualized on the edge of placental tissue and not within the villous structure.
Tissue clarification
Samples were washed four times in 2% donkey serum and PBS for 10 min at 21°C at 150 rpm prior to dehydration. Three 10-min washes of 50, 75 and 100% methanol were performed before immersing the tissue in the Visikol HISTO1 for 4 h at 21°C at 150 rpm. The tissue was placed in the Visikol HISTO2 solution overnight, prior to imaging the next day.

Image acquisition
Clarified samples were placed in a 12-mm Nunc™ Glass Bottom Dish (Thermofisher Scientific, TX, USA), and immersed in 500 µl of Visikol HISTO2 solution. Imaging was performed using a Leica SP5 AOB5 spectral confocal system in two-channel (Leica/ALEXA 488, Leica/ALEXA 633) optical sections at optimized intervals calculated by the imaging system (−1 µm per slice, 200–300 slices per sample) using a x20 PL APO NA 0.75 IMM CORR CS2 air objective. Optical detectors were set to 488–520 nm and 635–680 nm, respectively, to minimize crossover. To increase resolution, the confocal was set to 1024 × 1024 pixels, pinhole of 0.4 Airy, at 200 Hz acquisition speed, with line averaging. Negative controls (e.g., no primary antibody, or no secondary antibody) were used to adjust the confocal gain sensitivity to eliminate auto-fluorescence.

3D rendering
Imaged z-stacks were uploaded into Fiji-ImageJ as hyperstacks and the stacks were projected into either maximum intensity or 3D renderings.

RESULTS & DISCUSSION
We describe the developed protocol for the successful immunofluorescent labeling and clarification of the placenta of the NHP, which has been refined for confocal microscopy and quantitative analysis. We approached protocol development as an iterative process with the aim to maximize both reproducibility and the signal-to-noise ratio of our immunofluorescent labeling as well as to minimize the need for post-imaging processing. Our predetermined threshold for resolution was 3 µm; the terminal diameter of the capillary vessels. When visualized at 20x magnification (PL APO NA 0.75 AIR CORR CS2 objective), maximum-intensity projections of the clarified and immunolabeled NHP placental tissue allow for clear delineation of the capillary vessels from the syncytiotrophoblast tissue (Figure 2), and the 3D orientation of the placental micro-anatomy within the intervillous space (Figure 3). While other groups have used the Visikol HISTO solvent for placental clarification, our group found that those protocols were ineffective in the NHP placenta. Specifically, we found that antigen retrieval was essential, and multiple measures were required to decrease auto-fluorescence and maximize the intensity of the immunofluorescent labeling.

Experimental considerations

Auto-fluorescence
Fixed placental tissue, visualized with confocal microscopy in the absence of clarification or immunolabeling, has an abundance of natural auto-fluorescent noise (generated through laser-excitation of fluorophores in lipofuscin, elastin, collagen, heme byproducts and byproducts of cellular metabolism) making it unsuitable for quantitative analysis (Figure 4). Interfering auto-fluorescence can come from multiple sources including naturally occurring organic molecules and retained heme byproducts, and they can also be induced by the fixation process. While imaging techniques and post-imaging correction methods exist, these are unreliable and can reduce the desired signal from immunolabeled structures. For this protocol, duration of Zn Formalin exposure was minimized to decrease the amount of fixation-induced auto-fluorescence. Clearance of red blood cells can be achieved through saline perfusion of fresh placenta; however, this is technically difficult in the NHP and not always feasible. Rinsing the tissue with PBS followed by a H₂O₂-based bleaching solution under optimized conditions reliably removed speckling from retained heme byproducts. Lastly, minimizing heat exposure and exposing the tissue to LED photobleaching further decreased the background natural and fixation-induced auto-fluorescence.

Antigen retrieval
We found antigen retrieval to be necessary for our CD31 and CK7 antigens, with the temperature, duration and pH optimized as a balance between the two antibodies. In addition, numerous alternative primary antibodies and lectins were trialed under varying conditions (concentration, agitation, duration, temperature) and antigen retrieval steps (buffer type, pH, duration, temperature and pressure settings). Prolonged, high-
concentration washes under heat were required to obtain optimal signal strength and tissue penetration.

**Clarification & reversal**

Tissue clarification was carried out as per the manufacturer recommendations. While the use of the second solution (Visikol HISTO2) was listed as being optional for vascular tissues like the placenta, omitting this step dramatically decreased the effectiveness of the clarification process. Reversal of the clarification process and subsequent H&E-stained paraffin-embedded slides were generated to correlate our quantitative analysis with stereologic findings.

**Image acquisition & analysis**

ImageJ is a freeware program capable of rendering uploaded z-stacks into 3D and subsequently analyzing the geometry of the placental micro-anatomy. While filters exist for the smoothing of immunolabeled structures and subtracting out background noise, we sought to minimize the amount of post-imaging processing required prior to analysis by maximizing the signal-to-noise ratio of our immunofluorescent-labeling and clarification protocol. Prior to image analysis, the vascular channel must be isolated and made binary, a process whereby each voxel is analyzed and those that have an intensity below a specified threshold are removed while those that are above threshold are altered to a uniform intensity (Figure 5). This crucial step requires accurate determination of the set threshold, and z-stacks containing heme byproducts, unevenly stained structures, or increased auto-fluorescence result in the loss of labeled structures and the incorporation of unwanted noise interpreted as structures of interest. Through the refinement of our protocol, ImageJ becomes more sensitive and specific to our immunolabeled objects of interest and the subsequent quantitative analysis is more robust.

**CONCLUSIONS & FUTURE APPLICATION**

With the development of new analytic tools our ability to determine disease etiology and monitor response to therapy improves. The protocol as described in this paper allows for the generation of confocal imaged z-stacks with sufficient resolution to pursue reliable and reproducible quantitative analyses. By identifying and reliably quantifying micro-anatomic changes in highly-controlled, pathologic NHP pregnancies, precise mechanisms can be identified, explored and monitored in the absence of the inevitable confounding variables that complicate analysis of human pregnancies. When these methods are subsequently applied to human placenta studies, insights from NHP pregnancies can potentially translate to clinical breakthroughs and improvement in obstetric outcomes.

**FUTURE PERSPECTIVE**

As progress in the confocal imaging field continues to advance, the availability of commercially available products (such as Visikol HISTO) will allow for improved consistency in protocol reproducibility across multiple scientific disciplines and research institutes. For placental research, we anticipate that modification of the protocol described here could potentially be used for other 3D analyses including the localization of viral and bacterial antigens, cells of the immune system, membrane proteins and transporters, or genetic alterations (e.g., placental mosaicism).

**AUTHOR CONTRIBUTIONS**

JAS, VHJR and AEF conceived the study and experimental design. JAS performed the experiments with the assistance of JEG. JAS performed data acquisition. JAS wrote the manuscript with assistance from VHJR. All authors reviewed and approved the final manuscript.

**FINANCIAL & COMPETING INTERESTS DISCLOSURE**

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ETHICAL CONDUCT OF RESEARCH

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REFERENCES

Papers of special note have been highlighted as: of interest

1. Benirschke K. Pathology of the Human Placenta. Burton G, Baergen RN (Eds). Springer, Berlin, Germany, London, UK (2012).
2. Baergen RN. Manual of Pathology of the Human Placenta. Second Edition. Springer US, Boston, MA, USA (2011).
3. Funaro M, Ishida J, Aoki, I, Fukumizu A. Pathophysiology of placentation abnormalities in pregnancy-induced hypertension. Vasc. Health Risk Manage. 4(5), 1301–1313 (2008).
4. Kovt M, Schreiber L, Ellyashvili O, Ben-Harouah A, Abraham G, Bar. J. Pregnancy outcome and placental findings in pregnancies complicated by fetal growth restriction with and without pre-eclampsia. Reprod. Sci. 22(3), 316–321 (2015).
5. Retrospective review of 342 human pregnancies complicated by fetal growth restriction (72 with hypertensive disorders of pregnancy and 270 without). Investigators found high rates of placental vascular lesions (e.g., acute atherosis and mural hypertrophy) in both groups, and an increase in the number of villous lesions associated with maternal underperfusion (e.g., syncytial knots, villous agglutination, intervillous fibrin deposition and villous infarcts) in those pregnancies complicated by hypertensive disorders of pregnancy and growth restriction when compared with those pregnancies with isolated growth restriction.
6. Rainey A, Mayhew TM. Volumes and numbers of intervillous pores and villous domains in placentas associated with intrauterine growth restriction and/or pre-eclampsia. Placenta 31(7), 607–606 (2010).
7. Salafia CM, Vampolsky M, Misra DP et al. Placental surface shape, function, and effects of maternal and fetal vascular pathology. Placenta 31(11), 958–962 (2010).
8. Mayhew TM, Manwani R, Ohadice C, Wijesekara J, Baker PN. The placenta in pre-eclampsia and intrauterine growth restriction: studies on exchange surface areas, diffusion distances, and villous membrane diffusive conductances. Placenta 28(2–3), 233–238 (2007).
9. Peterson DA. Quantitative histology using confocal microscopy: implementation of unbiased stereology procedures. Methods 18(4), 493–507 (1999).
10. Mayhew TM, Ohadice C, Baker PN, Cropper IP, Mitchell C, Dang SS. Stereological investigation of placental morphology in pregnancies complicated by pre-eclampsia with and without intrauterine growth restriction. Placenta 24(2–3), 219–226 (2003).
11. Pittman Mayo R, Charnock-Jones DS, Burton GJ, Oyen ML. Three-dimensional modeling of human placental vascularization. Placenta 31(11), 958–962 (2010).
12. Haeussner E, Aschauer B, Burton GJ. Methods Mol. Med. 121, 371–392 (2006). Vascular corrosion casting of human placentas using vascular casting methods. When compared with normal term pregnancies, the fetal growth restriction group was found to have increased placental branching, decreased peripheral artery-to-vein ratios, and smaller vessel diameters.
13. Ribatti D. Vascular Morphogenesis Methods and Protocols. New York, NY, USA (2011).
14. Whiteley KJ, Pfarrer CD, Adamson SL. Vascular corrosion casting of the uteroplacental and fetoplacental vascular tree in mice. Methods Mol. Med. 121, 371–392 (2006).
15. Junaid TG, Bradley RS, Lewis RM, Aplin JD, Johnstone ED. Whole organ vascular casting and microCT examination of the human placental vascular tree reveals novel alterations associated with pregnancy disease. Sci. Rep. 7(1), 4144 (2017).
16. Yin G, Chen M, Li J et al. Vascular corrosion casting of normal and pre-eclamptic placentas. J. Thromb. Haemost. 14(6), 5535–5539 (2017).
17. Gong SP, Zhao YT, Yu YH. Vascular network modeling reveals significant differences in vascular morphology in growth-restricted placentas. Rev. Obstet. Gynecol. 4(3–4), 103–108 (2011).
18. Comparison of 20 placentas from human pregnancies complicated by fetal growth restriction and 20 control placentas using vascular casting methods. When compared with normal term pregnancies, the fetal growth restriction group was found to have increased placental branching, decreased peripheral artery-to-vein ratios, and smaller vessel diameters.
19. Sheppard C. Confocal Laser Scanning Microscopy. Shotton D (Ed.). Royal Microscopical, Oxford, UK; BIOS Scientific Publishers, NY, USA; Springer, Oxford, NY, UK, NY, USA (1997).
20. Price RL, Jerome WG. Basic Confocal Microscopy. Springer, NY, USA (2011).
21. Paddock DW. Confocal Microscopy Methods and Protocols. Springer, NY, USA, Imprint: Humana Press, NY, USA (2014).
22. Richardson DS, Lichtman JW. Clarifying tissue clearing. Cell 162(2), 246–257 (2015).
23. Merz G, Schwink V, Shah R et al. Three-dimensional rendering and analysis of immunolabeled, clarified human placental villous vascular networks. J. Vis. Exp. (123), e54348 (2017).
24. A video follow-up to the 2017 Placenta article by the same group that demonstrates the clarification and immunolabeling of a human placenta and begins the discussion of the potential quantitative analyses possible with these techniques.
25. Meier JG, Schwink V, Shah R, Aplin JD, Merz G. Clarification and 3-D visualization of immunolabeled human placenta villi. Placenta 53, 36–39 (2017). First reported use of Visiokol® HISTO™ in the placenta.
26. Richardson DS, Lichtman JW. Clarifying tissue clearing. Cell 162(2), 246–257 (2015).
27. Meier JG, Schwink V, Shah R, Aplin JD, Johnstone ED. Whole organ vascular casting and microCT examination of the human placental vascular tree reveals novel alterations associated with pregnancy disease. Sci. Rep. 7(1), 4144 (2017).
28. Marks PC, Preda M, Henderson T et al. Interactive 3D analysis of blood vessel trees and collateral vessel volumes in magnetic resonance angiograms in the mouse ischemic hindlimb model. Open Med. Imag. J. 7, 19–27 (2013).