Title
A new method for the sampling and preservation of placental specimens in low-resource settings for the identification of *P. falciparum* and analysis of nucleic acids

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Abstract
Collection, preservation, and shipment of histological specimens in low-resource settings is challenging. We present a novel method that achieved excellent preservation of placental specimens from rural Mali by using formalin fixation, ethanol dehydration, and long-term storage in a solar-powered freezer. Sample preservation success was 92%, permitting evaluation of current and past malaria infection, anemia, placental maturity, and inflammation. Using RNAscope® hybridization we were able to visualize cell-specific gene expression patterns in the formalin-fixed paraffin-embedded (FFPE) specimens. Additionally, our method entailed mirrored sampling from the two cut faces of a cotyledon, one for the FFPE workflows and the other for storage in RNAlater™ and RNA-seq.
Introduction

Neutral-buffered formalin is routinely used for tissue preservation including for the placenta [1–5]. For this project, we sought to preserve placentas to detect malaria infection. Formaldehyde can produce artifacts that resemble hemoglobin deposits, the waste product of the malaria parasite, particularly when acid formalin is used [6]. Even specimens fixed with neutral formalin can be subject to artifacts under conditions of prolonged fixation, elevated temperature, and high humidity [7]. We collected placental specimens in a rural location in Mali, West Africa, that is endemic for malaria (Plasmodium falciparum) and where ambient temperature often exceeds 40°C. We report a new collection protocol for placental specimens collected following delivery that minimized the formalin artifacts, required little on site infrastructure, and resulted in specimens that could be shipped on dry ice. Our collection and preservation protocol routinely produced high quality histological specimens suitable for the assessment of morphological features of the delivered placenta, including changes associated with active or past placental malaria infection, and suitable for in situ RNA hybridization. In parallel, we also collected specimens in RNAlater™ without fixation for molecular analysis of nucleic acids.

2.1. Collection and processing

The placentas came from women who participated in a longitudinal study on the Bandiagara Escarpment in Mali. The cohort and IRB approval is described in detail in Vincenz et al. [8, 9]. To obtain specimens from term placentas, we identified well-formed cotyledons within the central two-thirds of the maternal surface of the placenta. We then cut each cotyledon in half and dissected the histological specimen from the interior of one cut face and the specimen for nucleic acid analysis from the opposite face. The areas sampled consisted mainly of fetal villi and maternal intervillous space. We performed this mirrored sampling on two cotyledons per placenta generating 638 histological and nucleic acid specimens.

Histological specimens were placed in tissue cassettes and fixed with formaldehyde (Merck, purchased in Bamako, the capital city) freshly diluted 1:10 with phosphate buffered saline (PBS) to 3.7% final concentration. PBS was reconstituted using PBS tablets (Sigma-Aldrich) and distilled water. Fixation was for 36 hr. on ice followed by a 30 min. wash in 70% ethanol. The tissue cassettes were shaken off, transferred to plastic bags, and stored in a solar freezer (~ -20 °C) for up to 18 months prior to shipment on dry ice via World Courier to a -80 °C freezer at the University of Michigan. Specimens for nucleic acid analysis were processed as previously described [8].

2.2 Slide preparation and visualization

The tissue cassettes were removed from the -80 °C within a year of their arrival and submerged at room temperature in 70% ethanol prior to processing at the University's Tissue and Molecular Pathology (TMP) core. Tissue blocks were embedded in paraffin and two four-micron thick sections were cut from each sample and stained with hematoxylin and eosin (H&E) or Giemsa. Histology slides were examined under an
Olympus BX40 microscope. Polarized light along with location of pigments permitted hemozoin to be distinguished from formaldehyde artifacts [10]. The RNAscope® Multiplex Fluorescent V2 assay (Figure 2) was performed according to the user manual using unstained specimens [11]. RNAscope® slides were imaged on a Leica Stellaris confocal microscope.

We scored histological preservation as 0 for samples without fixation issues, as 1 for samples with damage limited to erythrocytes but still readable, and as 2 if the samples were completely unreadable.

Results & Discussion

Fig 1. Representative micrographs from specimens of malaria-infected term placentas collected in the field using our novel protocol. Arrows point to features of interest. A shows hemozoin, B shows gametocyte form, C shows trophozoite form, D shows schizont form. Scale bars are 20 μm.
The giemsa-stained specimens (Figure 1) show intact placental morphology, with excellent preservation of the villi, macrophages, neutrophils, and erythrocytes. Figure 1A shows brown hemozoin, the waste product of *P. falciparum*, located in perivillous fibrin of the placenta. Figure 1B shows the gametocyte stage of *P. falciparum* [12], which is the sexual form. Figure 1C shows a severe case of malaria with many *P. falciparum* trophozoites (activated feeding stage). Figure 1D shows a schizont, a mature stage of *P. falciparum*. The asexual stage parasites in A, C, and D indicate active infections. Panel B demonstrates the sexual erythrocytic stage of the parasite that is ingested by mosquitoes during a blood meal [12]. The presence of only hemozoin indicates past infection.

RNAscope® in-situ hybridization was performed to examine whether the preservation of RNA was sufficient to visualize the expression of specific genes. The RNAscope® probes were for two imprinted IncRNAs, KCNQ1OT1 and MEG3, and a protein-coding RNA, ERVW-1, which is a marker for trophoblasts. All three probes generated strong signals in different cells indicative of specific hybridization. Thus, the specimens collected with this protocol were of sufficient quality to provide spatial information on the expression of specific genes.

**Figure 2.** RNAscope® hybridization with probes for three genes. **A.** DAPI **B.** KCNQ1OT1 **C.** MEG3 **D.** ERVW-1 **E.** Merged. Scale bar is 20 μm.

We collected 638 histology samples from 322 placentas. We excluded 49 due to sample properties unrelated to fixation (e.g. ischemia). Of the remaining 589 samples, 544 (92%) samples were readable (score of 0 or 1). The 45 unreadable specimens could not be assessed for malaria infection due to loss of structure reminiscent of freezing artifacts. We did not attempt to use 30% sucrose as a cryoprotectant due to the inability to assess osmolarity artifacts in the field.

The protocol presented here will improve the collection of placental specimens in low resource settings. Formaldehyde is widely available, but due to its toxicity, appropriate training of personnel is essential. Solar freezers are available in most urban areas and can be installed in rural settings. Maintenance costs consist mainly in the periodic exchange of lead acid batteries. The overall quality of the FFPE specimens was excellent and permitted the evaluation of current and past malaria infection, anemia,
placental maturity, and inflammation, as well as visualization of gene expression using RNAScope®. The RNA specimens produced high quality data using targeted RNA-seq [13].
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