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TITLE: AI-Assisted 3D Ultrasound for Rapid Diagnosis of Ocular Trauma Injuries

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CONTRACTING ORGANIZATION: Case Western Reserve University, Cleveland, OH

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ing; thus, limiting the utility of optic based methods such as direct ophthalmoscopy or OCT. Small foreign bodies, may not be well visible in CT imaging, even at highest resolutions. Ultrasound has been used for diagnosis and treatment of ocular injuries. Conventional 2D ultrasound requires skilled users and are subject to large variability. We propose to develop a novel 3D ultrasound system to greatly facilitate diagnosis and treatment of ocular injuries. Our gentle-touch approach via single sweep scanning minimizes the risk of additional injuries. We have created a prototype of the system resulting in anatomically correct volumes of ex vivo eyes. Initial deep learning results show promise in automated analysis of 3D data.

15. SUBJECT TERMS

3D, Ultrasound, Ocular injuries, trauma, retinal detachment, intraocular foreign bodies

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1. Introduction

The overarching goal of this project is to create a 3D ultrasound system for imaging of the eye to remove or minimizes problems associated with current ultrasound imaging of the injured eyes, especially among service members impacted by military trauma. Our approach is to build a prototype 3D ultrasound system and test its various aspects using in vitro, ex vivo and in vivo preclinical models of ocular injuries. Our plan is to test our system on realistic models of trauma, especially those in military including retinal detachment, foreign bodies, variation of optic and vascular integrity. We utilize computer modeling and physical experiments to develop advanced image acquisition, volume reconstruction and enhancement to eventually validate on a rabbit model of ocular injuries. Where applicable, to better understand progress, we included the related major tasks as proposed in our statement of work (SOW).

2. Keywords

3D, ultrasound, ocular injuries, military trauma, retinal detachment, optic nerve sheath diameter, imaging, diagnosis, intraocular foreign bodies, deep learing.

3. Accomplishments

The overall goal of this project is to

- Develop and optimize a 3D ultrasound system
- Experimentally validate on ex vivo and in vivo models of ocular injuries

To achieve these goals, the following accomplishments are reported based on the statement of work:

- IACUC obtained

We obtained IACUC approval for our animal study from Case Western Reserve University to conduct the research proposed in this project.

Date accomplished: July 2021.

- Acuro obtained

We obtained ACRU approval for our animal study. We both IACUC and ACURO approval in hand, we are set to start the preclinical studies our project as outlined in the statement of work. Date accomplished: November 2021.

Major Task 1:

Developing 3D imaging

We purchased and obtained an ultrahigh frequency wireless portable ultrasound probe from Clarius (Clarius inc, Vancuver, CA) to conduct experiments.

Assembly with stabilizer and initial tests:

We have created the first generation of our 3D ultrasound system using a Clarius ultrahigh frequency probe. The system is currently able to obtain 3D volumetric data using a fully controlled software-hardware integration via tools pro-

vided by Clairus as well as our custom-made units. Our 3D system houses the Clarius probe in a custom-made 3D printed holder which can adapt to a variety of connectors for ex vivo and eventually in vivo imaging. Fig. 1 shows our current system in action during an experiment. Fig. 2 shows an example of volume reconstruction using an ex vivo porcine eye. Currently, the motion mechanism is provided us-

ing a precise stepper motor instead of using a camera stabilizer. This was mainly due to speed up prototyping using our past experiences with this setup and starting the ex vivo experiments.

Percentage accomplished: 80%.

Delay in accomplishing this part was partly due to finding a graduate student and partly related to acquiring the required tools.

Future works: We will optimize intensity equalization and reduction of artifacts, mostly related ex vivo testing (e.g. reflec-

tion from phantom boundaries, tissue integrity during shipment). We will continue to investigate the

suitability of a gimbal camera stabilizer and use our current setting as the benchmarked operation.

Fine tuning and validation on phantoms

We found empirical parameters to create reproducible isovoxel image volumes. We developed ultrasound fine targets using gelatin and suture wires with diameter < 30mu and created 3D volumes which showed consistent results. Examples of wire target image is shown in Fig. 3.

Date Accomplished: April 2022.

Major Task 2:

Developing iterative enhancement methods:



Figure 1: 3D ultrasound prototype during an ex vivo testing



Figure 2: Multiplanar 3D visualization of an ex vivo porcine eye acquired by our prototype



Figure 3: 3D PSF estimated from fine wire target experiment

PSF estimation: We have conducted precision experiments on custom-made fine target phantoms to fully evaluate imaging characteristics of the developed 3D ultrasound. These included determining 3D point-spread-function (PSF), scanning speed requirement, data acquisition characteristics including number of frames, voxel size etc. Fig. 3 shows volumetric PSF in two perpendicular views to be used in inversion deblurring methods.

Percentage accomplished: 80%.

Future work: We continue to work on the PSF inversion algorithms for image improvement to include ansiometry of the PSF.

Scan conversion and Deconvolution: Since we are using the standard data from the commercial probe, no scan conversion was found to be required. Deconvolution was achieved using Richardson-Lucy (RL) with spatially variant PSF as obtained in the previous sub-task.

Percentage accomplished: 80%.

Future works: Once our experimental dataset become suitable for training a deep learning, we will convert the time-consuming RL deconvolutions to faster implementation.









Motion corrected slice

Figure 4: Demonstration of our motion compensation algorithm performed on data from ex vivo testing. Only a single slice of 3D volume is shown

introduced within [-5 5] pixels

Motion compensation: Mechanical motion of the probe may create slight jitters due to damped infinitesimal relative probe-tissue relative positions. During in vivo, physiological motions, e.g. respiration, pulsation can create motions that will induce misregistration across adjacent planes in a 3D acquisition. We developed methods based on frame-to-frame similarity to remove such artifacts for enhanced volume acquisition. Fig. 4 demonstrates performance on a porcine model of ocular foreign bodies where artificial motions and added noise were significantly removed by our algorithms.

Percentage accomplished: 70%.

Future work: Our algorithms work well in ex vivo models with small motions. We will continue to develop for in vivo and potentially larger motions.

Testing on standard small part and phantoms and ex vivo porcine eyes:

We used our developed 3D US prototype to acquire image volumes from wire phantoms (Fig 3) and ex vivo porcine eyes. Fig 5 shows a typical experimental setting where ex vivo porcine eyes are embedded in tissue mimicking gelatin phantoms and scanned using our 3D system prototype. The model allows more reproducibility and creates the ability to image from multiple angles to verify volume reconstruction performance and rule out orientation-dependent artifacts. This model will also allow better cross modality registration as we plan to perform CT imag-



Figure 5: Ex vivo porcine eye case in gelatin for testing

ing of these ex vivo models of ocular injuries such as IOFBs and retinal detachment. Date accomplished: July 2022.

Major Task 3:

Developing deep learning for automated segmentation and identification:

Ex vivo porcine eye experiments for training GAN: We developed deep learning methods for automatic segmentation of the ciliary processes in the anterior segment. Though these methods are currently applicable to the anterior segment, mainly due to the nature of previously collected dataset, they can be easily extended to whole eye imaging to address problems associated with traumatic injuries as

proposed in our grant proposal. Fig. 6 shows our deep learning segmentation of the ciliary body (prediction) as compared to delineation performed by an expert.

Manual segmentation of ex vivo 3D US: To prepare the developed GAN for our tasks (IOFB and RD detection) we continue to perform ex vivo experiments and use manual segmentation as the ground truth.

Percentage accomplished: 60%.

Future work: We have developed deep learning segmentation methods using a previously acquired dataset by our group. This dataset is limited to the anterior segment. We have already created models for successful induction of IOFBs and RDs in ex vivo eyes. We will



Figure 6: Deep learning segmentation of ocular structure (ciliary body) using our developed methods (prediction) compared to manual segmentation

continue to build on this model to cover the whole eye and include numerous instances of ocular injuries, IOFBs and RD with manual segmentation for training our deep learning model.

Major Task 4:

Ex vivo validation studies on porcine eyes

Part of the ex vivo validation studies is related to ongoing ex vivo testing. While so far, the focus has been on accomplishing the 3D US system performance, we have initiated parallel validation studies using CT. The focus of our work in the next few months will be on these validation studies, which will in turn prepare us for in vivo testing.

Percentage accomplished: 30%.

Future work: we continue our ex vivo testing for both developing deep learning algorithms as well as preparing samples for cross-modality validation as indicated in Major task 4. Some microCT studies have been already performed and plans are developed for ex vivo CT and Cryo imaging in the next following months.

Evaluation of images and analyzing data:

Anticipated date: April 2023

Major Task 5:

In vivo studies on rabbits Anticipated date: June 2023 Manual segmentation of in vivo 3D US Anticipated date: June 2023

What opportunities for training and professional development has the project provided?

This project engaged a full time PhD student who is dedicating his PhD work on ultrasound imaging of the eye. He is gaining and developing skills in both system development and artificial intelligence via developing novel deep learning analysis methods.

- How were the results disseminated to communities of interest?

So far we have presented our works in technical conferences included SPIE (past) and IUS IEEE (upcoming) and have published one manuscript Translational Vision Science & Technology journal

- What do you plan to do during the next reporting period to accomplish the goals?

Plans for future works are explained for each milestone above. Overall, once we achieve our goals in reliability and accuracy of the developed system, we plan to initiate our in vivo studies to test the utility in more realistic situations. As part of challenges, we anticipate a learning process for inducing relevant injuries similar to those seen in military trauma in rabbit eyes. Our ex vivo studies are meant to equip us with the required skillset to be better prepared for these challenges.

4. Impact

Our goal is to fill a crucial gap in facilitating ocular imaging in vision threatening situations where no other modalities are applicable. Important applications include: identification of IOFBs, surgical planning, and intraoperative guidance, diagnosis of retinal detachment, evaluation of optic nerve as a surrogate of increased intracranial pressure and checking vascular integrity to assess hemorrhage and bleeding. Ultrasound is currently used in treatment of traumatic ocular injuries, especially among service members. However due to 2D planar imaging and requiring manual scanning is risky on injured eyes.

What was the impact on the development of the principal discipline(s) of the project?

Our method will greatly improve volumetric acquisition which in turn allows automated analysis, for example automated detection and segmentation of IOFBS. We anticipate a future portable product that can be easily implemented by non-skilled users in remote and austere environments such as low-level military triage or in emergency department for in ocular injuries in general public.

What was the impact on other disciplines?

The core system development and algorithms developed in this project can be potentially extended to 3D ultrasound imaging of other organs.

What was the impact on technology transfer?

Demonstrating clinical utility in ocular trauma can help us effort in attracting industrial partnership to translate this research into a commercial product for 3D imaging of the eye.

What was the impact on society beyond science and technology?

Nothing to report during this reporting period.

5. Changes/Problems

Changes in approach and reasons for change

The initial idea for creating 3D ultrasound was to use a professional camera gimble with accurate rotation capability to create multi-plane sector scanning. While we are searching for a good hardware for this purpose, we found it helpful to use small size OEM linear motors and performing linear scanning. The acquired volumes are quite satisfactory and will be used as baseline for any other alternative approach including sector scanning. No other problem or changes occurred or seen in our project so far.

Actual or anticipated problems or delays and actions or plans to resolve them

A delay was related to recruiting a PhD student as explained in this report. No other delays or problems to report.

Changes that had a significant impact on expenditures

Nothing to report.

Significant changes in use or care of human subjects, vertebrate animals, biohazards, and/or select agents

Nothing to report.

Significant changes in use or care of human subjects

Nothing to report.

Significant changes in use or care of vertebrate animals

Nothing to report.

Significant changes in use of biohazards and/or select agents

Nothing to report.

6. Products

Publications, conference papers, and presentations Within this reporting period, we presented one conference paper at SPIE

Ahmed Tahseen Minhaz, et al., Deep learning segmentation of ciliary tissues using 3D ultrasound biomicroscopy (3D-UBM) images, Medical Imaging 2022: Ultrasonic Imaging and Tomography, vol. 12038, Pages 233-237, 2022

A conference paper is accepted for presentation at 2022 IEEE IUS and will be included during the next reporting period.

Journal publications.

We have also submitted a manuscript to Translational Vision Science & Technology

Ahmed Tahseen Minhaz, et al., Deep learning segmentation, visualization, and automated 3D assess-

ment of ciliary body in 3D ultrasound biomicroscopy (3D-UBM) images, in review

Books or other non-periodical, one-time publications. Nothing to report.

Other publications, conference papers and presentations. Nothing to report.

Website(s) or other Internet site(s) Nothing to report.

Technologies or techniques Nothing to report.

Inventions, patent applications, and/or licenses Nothing to report.

Other Products

Nothing to report.

7. Participants & Other Collaborating Organizations

What individuals have worked on the project?

Name: Mahdi Bayat, PhD

Project Role: PI

Nearest person month worked: 6

Contribution to Project: Dr. Bayat managed and directed this project and supervised graduate students. He also contributed in developing algorithms, collecting data, revising manuscripts and holding regular weekly meetings.

Name: Ahmed Tahseen Minhaz

Project Role: PhD Student

Nearest person month worked: 8

Contribution to Project: Tahseen worked on several aspects of this project. He developed a 3D ultrasound prototype and tested on several phantoms he made. He participated in group meetings and created abstracts and manuscripts. He also developed deep learning methods for segmentation of ocular tissues, image improvement and microvascular imaging. Name: David Wilson, PhD Project Role: Co-I Nearest person month worked: 1

Dr. Wilson participated in group meetings and provided supervision and guidance, especially in developing 3D image analysis algorithms. He also reviewed manuscripts and reports and provided valuable feedback.

Name: Faruk Orge, MD Project Role: Co-I Nearest person month worked: 1

Dr. Orge participated in group meetings and provided supervision and guidance, especially in practical and clinical aspects of the project including the development of ex vivo and preclinical models of traumatic ocular injuries. He also reviewed manuscripts and reports and provided valuable feedback.

Has there been a change in the active other support of the PD/PI(s) or senior/key personnel since the last reporting period?

Mahdi Bayat (PI): Nothing to Report.

Faruk Orge (Co-I): Nothing to Report.

David Wilson (Co-I):

New Active

Title: 3D Microscopy with Ultraviolet Surface Excitation (3D-MUSE)

Major Goals: The goal is to create a new section-and-image microscopy approach. Status of Support: Active Project Number: R01 EB028635 Name of PD/PI: Wilson Source of Support: NIH Primary Place of Performance: Case Western Reserve University, Cleveland, OH Project/Proposal Start and End Date: (MM/YYYY) (if available): 07/2019-03/31/2023 Total Award Amount (including Indirect Costs): Overlap: none

Title: System-independent quantitative cardiac CT perfusion

Major Goals: We will develop quantitative methods for accurate assessment of blood flow perfusion in myocardial tissue, enabling determination of micro-vascular disease. This project requires image registration, deep learning image segmentation, and parameter estimation.

Status of Support: Active Project Number: R44 HL156811 Name of PD/PI: Wilson Source of Support: NIH Primary Place of Performance: Case Western Reserve University, Cleveland, OH Project/Proposal Start and End Date: (MM/YYYY) (if available): 09/2021-08/2023 Total Award Amount (including Indirect Costs): Overlap: none

Title: Mapping the Autonomic Pathways in 100 Human Vagus Nerves in a Demographically Representative Sample (MAP-100VN)

Major Goals: The vagus nerves connect the brainstem to most organs in the torso—including the heart, lungs, pancreas, spleen, stomach, and intestines—to control and sense our body's inner workings. Electrical stimulation of the vagus nerve has been used to treat epilepsy, rheumatoid arthritis, and heart failure, among many other conditions; however, insufficient anatomical data are available to map the ~100,000 neurons of the human vagus nerve to their target organs for development of improved therapies, with increased efficacy and decreased side effects. Therefore, we will use our team's integrated and comprehensive expertise to characterize 100 human vagus nerves using different imaging methods, conducting the most comprehensive imaging and mapping of the vagus nerve to date, spanning gross anatomy through the body to micron-scale structure of individual neurons, to seed and accelerate the development of new neuromodulation therapies for treatment of disease.

Status of Support: Active

Project Number: CWRU NIH SPARC 75N98022

Name of PD/PI: Shoffstall. Wilson is a co-I.

Source of Support: NIH SPARC REVA

Primary Place of Performance: Case Western Reserve University

Project/Proposal Start and End Date: (MM/YYYY) (if available): 09/21/2022 - 09/20/2025

Total Award Amount (including Indirect Costs):

Overlap: none

Closed Grants

Title: Cancer Imaging and Therapy Analysis Platform (CITAP).

R44 CA213601 (mPI: Gargesha, BioInVision, and Wilson) 09/20/16 - 08/31/20. annual sub-contract to lab. We will develop cryo-imaging and software for analysis of cancer.

Title: Interdisciplinary Biomedical Imaging Training Program.

NIH T32EB007509. PI: Wilson. 9/1/2007-8/31/2022. This training grant supports predoctoral trainees in Biomedical Engineering and Physics in imaging.

Title: Image-guided EMT inhibition for treating metastatic breast cancer.

(NIH R01CA194518) PI: Lu, co-I: Wilson. 7/1/2015-6/30/2020. This project will be focused on developing new Image-guided therapeutic strategy by targeting proteins in key biological events associated with metastasis to treat life-threatening metastatic breast cancer. Dr. Wilson's role is image analysis.

Title: Three-dimensional ultrasound imaging for ophthalmology.

Case-Coulter Translational Research Partnership. 9/1/2018-8/31/2019. mPIs: Wilson and Orge.

What other organizations were involved as partners?

Nothing to report.

8. Special Reporting Requirements

See attachment.

9. Appendices

None.