A game for crowdsourcing the segmentation of BigBrain data

Arno Klein ‡
‡ Sage Bionetworks, Seattle, United States of America

Executive summary

The BigBrain, a high-resolution 3-D model of a human brain at nearly cellular resolution, is the best brain imaging data set in the world to establish a canonical space at both microscopic and macroscopic resolutions. However, for the cell-stained microstructural data to be truly useful, it needs to be segmented into cytoarchitectonic regions, a challenge no single lab could undertake. The principal aim of this proposal is to crowdsource the segmentation of cytoarchitectonic regions by means of a computer game, to transform an arduous, isolated task performed by experts into an engaging, collective activity of non-experts.

Keywords

BigBrain, brain imaging, histology, microstructure, neurons, crowdsourcing, game, gamify

Research & Related Other Project Information

Project Decsription

The principal aim of this proposal is to crowdsource the segmentation of brain histological data, specifically the cytoarchitectonic regions of the hippocampi of the human brain, by
means of a computer game. Currently, only human experts perform reliable cytoarchitectonic labeling at a very slow and small scale, whereas we propose to enlist many human non-experts to engage in a distributed version of this task at a quick and large scale. By turning this arduous, isolated task into an engaging, collective activity, we hope to radically change the way anatomists approach segmentation/labeling.

To support this aim, we must (1) prepare expert (gold standard) labels to a subset of the hippocampal sections to evaluate crowdsourced results, and (2) aggregate the crowdsourced results to label the hippocampi. For our first exploratory aim, will explore how our approach generalizes to all other brain regions, and for our second exploratory aim, we will train a supervised learning algorithm on the crowdsourced results and evaluate how closely the automated approach matches human assessments.

We will use the BigBrain, a high-resolution 3-D model of a human brain at nearly cellular resolution ($20\mu m$ isotropic) based on reconstruction of 7,404 histological sections stained for cell bodies. The BigBrain is extremely important to the neuroscience community because it represents whole-brain histological data with accompanying magnetic resonance imaging data. It is the best brain imaging data set in the world to establish a canonical space at both microscopic and macroscopic resolutions. However, for the cell-stained microstructural data to be truly useful, it needs to be segmented into cytoarchitectonic regions, which makes it the perfect focus for this project. Labeling of the BigBrain will establish an ex vivo atlas, a common space for neuroimaging data whose labels will provide a consistent, convenient, and meaningful way to communicate, classify, and analyze biomedical research data set in that space.

**Facilities & Other Resources**

**Sage Bionetworks**

**Laboratory:** N/A

**Clinical:** N/A

**Animal:** N/A

**High Performance Computing Resources:** Sage Bionetworks uses a combination of scalable cloud-based storage and analytical computational resources and its own computational facilities. The cloud-based services are procured from Amazon Web services on a fee for service basis and provide a cost-effective solution to variable needs, technology upgrades and support. Sage Bionetworks develops and operates two software as a service platforms, Bridge and Synapse, as resources for the broader scientific community. Both these systems operate on cloud-based infrastructure. Internal research projects also have access to the Sage Bionetworks high performance computing cluster, maintained through a partnership agreement with the University of Miami.

Additional servers used by Sage scientific staff are co-located at the Fred Hutchinson Cancer Research Center computing facilities. All networked file systems, databases, and
home directories are backed up using Veritas software to a robotic tape library. Tapes are taken off-site each month for disaster recovery.

**Additional facilities/resources:**

**Sage Bionetworks Bridge platform** is designed to support the design and execution of adaptive clinical trials delivered through smartphone platforms. Participants in a study interact with the Bridge server through an app which is custom-designed for each particular study. The Bridge server provides a way to configure a study with a set of survey questions to ask study participants and way to schedule times for the various survey and app data collection tasks to run. Storing this configuration on the server allows researchers to dynamically adjust the study as data is collected without distributing new builds of the app to participants.

The Bridge server provides services for study participants to create an account, authenticate, and manage informed consent to participate in a research study. The app will periodically save survey and sensor measurements to the server; client-side SDKs help manage transmission of his data over intermittent mobile network connections. Study data is stored separately from readily identifying user account data so that study participants can be deidentified and data shared protecting their anonymity. Once deidentified, study data will be stored on Sage Bionetworks Synapse platform.

**Sage Bionetworks Synapse platform** will be used to support the complex, interrelated analyses described in this project. Synapse is an informatics platform for collaborative, data-driven science that combines the power of community-based modeling and analysis with broad access to large datasets, which will enable the development of more predictive computational models of disease. Synapse is built as a web service-based architecture in which a common set of services is accessed via different sets of client applications, including a web portal and integrations with multiple analysis environments.

Synapse is designed to allow users to bundle together and publish the relationships, annotations, and descriptions of files that may live in multiple locations. This may be files stored in Synapse own native storage location (Amazon S3), data living on local file systems, or code from GitHub. As long as the storage location is accessible via http/ftp it can be accessed through Synapse. Synapse includes analysis clients for users working in the R and Python programming languages, and a tool that runs at the program line of the Linux shell, providing basic functionality to interact with the system no matter what analysis tools the analyst would like to use. These tools allow users to query and load data, post results, and create provenance records directly from the command line. The output of any analysis can be stored in the Synapse Amazon S3, or externally and indexed in Synapse just as the underlying data. The Synapse web portal is an environment for sharing data, results, methods and tools, and is a place that enables the tracking of analysis steps and publication of analysis results to collaborators and eventually the broader community. The Synapse web portal will allow researchers to publish analysis results and track project information. A Provenance visualization tool will allow users to formally track the
relationship among resources in the system and better document and communicate each analysis step.

The Synapse system is operated as a hosted service offering from Sage Bionetworks, requiring no installation of software or IT burden on the collaborating institutions. Researchers will be able to create accounts in the system and immediately use the system to collaborate.

At this time Synapse is hosted at http://synapse.org, and is being used to host a variety of bio-molecular data sets and analytical pipelines to curate this data. Sage Bionetworks has successfully used Synapse to support a number of large scale collaborative projects including open challenges in the predictive modeling of breast cancer survival and the TCG A Pan-Cancer working group.

Scientific Environment: Sage Bionetworks leases approximately 3,900 sq ft of office space on the Fred Hutchinson Cancer Research Center campus (FHCRC or the Center.) As part of the agreement with FHCRC, all Sage Bionetworks staff have full access to the Center’s research. Sage Bionetworks has a services agreement with FHCRC for facilities related logistics.

Centre de Recherches Interdisciplinaires (CRI)

Clinical: N/A

Animal: N/A

High Performance Computing Resources: N/A

Laboratory: Inserm U1001 lab in the Faculty of Medicine of the Paris Descartes University, working on Robustness and evolvability of life.

The lab offers the following facilities:

Microscopy:
- Axiovert 200 M and Axioplan 2 and Observer Z1 from Zeiss
- Eclipse Ti, Eclipse E200 from Nikon
- MetaMorph (software)

Microfluidics:
- SEM
- Chemostat

llumatool tunable lighting system
Microplates reader:
- iEMS (Labsystems)
- Victor3 (Perkin Elmer)
- Infinite (tecan)

Research in this lab, headed by F. Taddei and A. Lindner, is focused on studies into key issues such as aging on one hand and the origin of cooperation on the other, using state-of-the-art systems and synthetic biology approaches, including computer modelling, wet-lab experiments, robotics and nano-fabrication methods.

**Additional facilities/resources**:

**The MOOC Factory**

A platform to create MOOCs to support the CRI’s own education programmes but also to disseminate its interdisciplinary approaches to a larger public. In the factory, online courses focused on practice and experimentation are developed by innovators who wish to create courses for the general public, with a special emphasis for teachers, parents and children.

**OpenLab (DIY research FabLab)**

A place where students and entrepreneurs can prototype, code, hack and take part to the creation of connected objects and captors, around a multidisciplinary team.

The Lab has all the basic tools to prototype rapidly captors and interface them with web apps. Students can also use 3D printers, a laser cutter and a digital milling machine. A DIY-bio lab is also hosted at Cochin.

The OpenLab is a key resource to build and nurture an extended and diverse CRI community of bright minds. In less than two years, the OpenLab has attracted more than a hundred candidates and has selected 15 promising teams of young entrepreneurs on topics as diverse as quantified self, green mobility or open health.


**Education experience**

The CRI offers an Interdisciplinary curriculum, which spans the range from undergraduate (one Bachelor programme: Frontières du vivant) through masters (2 programmes: Interdisciplinary Approaches to Life Sciences and EdTech) to PhD studies in the fields of interdisciplinary life sciences and learning sciences at Sorbonne Paris Cité, one of France’s major university consortiums bringing together four Paris universities and four higher education and research institutes. The CRI hosts programmes for about 70 undergraduate students and 170 graduate students per year, in around fifty associated laboratories in the best French and international institutions, including the Rockefeller Institute, EMBL, Peking
University, Harvard University, and Imperial College London. The academic programmes are supervised and evaluated by an international scientific committee.

**Scientific Environment:**

The CRI leases approximately 3,000 square meters in the heart of Paris, in Le Marais. These dedicated facilities host visiting professors, a variety of courses, and many student clubs. They are complemented by 350 square meters of labs in the Institut Cochin.

The CRI organizes special programmes like the Leadership programme, which brings top-notch young researchers in Life Sciences to develop and implement innovative educational projects. During the three days of this programme, everybody present at the CRI can take part and interfere with the selected researchers.

The CRI also hosts Scientific Clubs created by students on a wide range of subjects of their choice such as interdisciplinarity, innovation, social and environmental development, science and society... The clubs give students the opportunity to get involved in an extracurricular activity that matches their interests and adds creativity, collaborativity and design thinking to their skill set. Current active clubs include WAX (aiming at creating and developing innovative and collaborative tools to make science fun, accessible to all and useful), Fabelier (a lab to create new things with/for the web) and Gamelier (for those who want to meet up to make games together, with a focus on educational and scientific games).

iGEM (international genetically engineered machine): the first French synthetic biology team for the MIT-sponsored iGEM competition. In this club, the CRI students have collected many awards through the years. In 2013, they were honoured with the Grand Prize at the World Championship Jamboree.

All the CRI activities, whether they are primarily focused on Education, Research or Entrepreneurship, attract and build up an exceptional community of bright minds inhabited by a common vision that knowledge sharing and open collaboration are important keys to our future. The diversity, quality and benevolence of the ever growing CRI community makes this place a unique resource to forge serendipity and unlock the unexpected.

The CRI hosts the Liliane Bettencourt Interdisciplinary Programme, which spans the range from undergraduate through masters to PhD studies. This 200-strong group comprises students from all the sciences, who carry out research in the best French and international institutions, including the Rockefeller Institute, EMBL, Peking University, Harvard University, and Imperial College London. The academic programmes are supervised and evaluated by an international scientific committee. CRI has initiated projects such as the MIT award-winning Paris synthetic biology iGEM team and interdisciplinary approaches to web sciences, Fablab hacker space. Researchers at the CRI are involved in projects at the interface of science, education and society, ranging from the Paris Montagne Science Festival for young children to outreach programmes for high school students from disadvantaged environment (Science Ac’87) and graduate students. CRI organizes Learning through Research workshops for students and lecturers across the world.
(Europe, China, Indonesia) and has launched the WISER-U as an international platform for students and scientists to design and develop new technologies for sharing ideas and creating projects together. Research at the CRI “Evolutionary Systems Biology” INSERM team, headed by F. Taddei and A. Lindner is focused on studies into key issues such as Aging on one hand and the origin of Cooperation on the other, using state-of-the-art systems and synthetic biology approaches, including computer modelling, wet-lab experiments, robotics and nano-fabrication methods.

The CRI’s dedicated facilities host visiting professors, a variety of courses, and many student clubs. It is host to Laboratory facilities of the U1001 unit of the French National Institute of Health & Medical Research (INSERM) in Paris-Descartes University’s Medical School; The OpenLab, a prototyping workplace and projects accelerator created to lower the entry barriers of technological and social innovation for students, scientists, designers, engineers, and entrepreneurs alike; The Game Lab, a place for students and researchers to learn about game creation and to develop their scientific games; The MOOC (Massive Open Online Courses) Factory, where online courses focused on practice and experimentation are developed by innovators who wish to create courses for the general public, with a special emphasis for teachers, parents and children.

**McConnell Brain Imaging Center (BCI) at McGill University**

**About the Center**

The McConnell Brain Imaging Centre (“The BIC”) is a multidisciplinary research centre dedicated to advancing our understanding of brain functions and dysfunctions and the treatment of neurological diseases with imaging methods. It is also one of the largest, multimodal brain imaging service platforms worldwide (largest in Canada), serving a community of 100+ investigators and generating a volume of more than 3,500 research scans per year in 7 imaging cores: MRI, PET, SPECT, MEG, EEG, etc.

Our Centre is also renowned for developing and distributing software applications and for having created multiple spin-off biomedical imaging companies.

**Core Facilities**

The unique platform infrastructure of the McConnell BIC is available and accessible to all researchers.

We offer one of the largest brain-imaging research and service platform worldwide. All imaging cores are hosted under the unique roof of the Montreal Neurological Institute (MNI) and hospital. Every year, the McConnell BIC scans >3,500 research participants with a multimodality of 7 research-dedicated scanners for humans and animal models.

**University of Pennsylvania**

Penn is home to a diverse body of over 20,000 students and over 4,000 faculty in its 12 leading graduate and professional schools. Penn’s schools are located on a compact
campus, the geographical unity of which supports and fosters its multidisciplinary approach to education, scholarship, and research. Research and research training are substantial and esteemed enterprises, bolstered by an annual University budget of $6 billion. Penn’s 165 research centers and institutes bring together researchers from multiple departments, schools, and disciplines, and interdisciplinary collaboration is a key theme for Penn’s academic enterprises.

Penn and the Perelman School of Medicine provide a truly outstanding intellectual environment in which to conduct the research we propose. The School of Medicine, and the Department of Radiology in particular, are consistently among the top five recipients of biomedical research funding in the nation. Virtually any kind of biomedical research is represented at Penn, and few other institutions offer a comparable breadth and depth of expertise and resources. This includes a wide range of imaging resources and expertise, including MRI, CT, 3D tomosynthesis, PET, 3D ultrasound, histology, and confocal microscopy.

An ongoing initiative to consolidate research in neuroimaging and brain function on the Penn campus is entering its second phase, and will result in nearly all neuroimaging investigators being located in newly renovated space centrally located on campus. By the end of 2015, Dr. Yushkevich’s laboratory will be located in the Richards and Goddard Laboratories Buildings, a designated National Historic Landmark designed by the renowned architect Louis Kahn to facilitate research interactions. Although the Penn campus is already very compact, consolidating the many faculty members involved in neuroimaging research will facilitate broader participation in projects, seminars, workshops, and interdisciplinary training, and will provide an outstanding environment for trainees to interact with faculty and with each other.

The primary site for this project will be the Penn Image Computing and Science Laboratory (PICSL) in the Department of Radiology. PICSL has an established track record of developing novel methodology for biomedical image analysis, disseminating methodology through open-source software, and applying it to relevant clinical problems through strong collaborations with clinical researchers at Penn and beyond. PICSL is home to over 20 graduate students, postdoctoral fellows and research staff. In summer 2015, PICSL will move to occupy a whole floor in the D tower of the Richards building, and will be steps away from the laboratories of leading neurologists and neuroscientists.

**Computing Resources**

**Computer Hardware and Software Resources**

The computing resources for this project will be available through the Neuroscience Neuroimaging Center (NNC), a NINDS P30 Institutional Center Core. John Detre is the PI of the NNC, and Paul Yushkevich is the PI of the Neuroinformatics Core, which maintains the NNC computing resources. The NNC maintains a state-of-the-art data processing facility with distributed computing, data storage and archiving, integration of neuroimaging
data analysis environments, and provides system administration support. The NNC high-performance computing cluster consists of

- 576 dedicated compute cores, from 21 compute nodes with dual 8-core Intel Xeon E5-2450 2.10GHz CPUs and 64GB RAM, and 30 compute nodes with dual 4-core Intel Xeon 2.83GHz E5-440 CPUs with 16GB RAM
- A dedicated 16-core head node (running Rocks and SGE) manages cluster operations
- Three dedicated file servers managing high-speed (6GB/s and 4GB/s) RAID-6 devices for over 200TB of formatted storage.
- A high-speed 10GbE internal cluster network and a Gigabit external network connection.
- A dedicated tape backup system with 100TB capacity.

The cluster is located in a University-run commercial-grade server room with redundant power supplies; UPS power backup systems; fire suppression system; and 24-hour restricted access and security.

Additionally, each investigator will have a personal workstation (Apple iMac or similar) with gigabit connection to the University of Pennsylvania Health System (UPHS) intranet, as well as fast Internet access. A range of software is available on the investigators' workstations and on the HPC cluster, including multi-platform software development tools, scientific computing packages, functional and structural MRI analysis frameworks; statistical packages, word processing and presentation applications, etc.

Resources for Open-Source Software Development and Dissemination

PICSL has recognized expertise and a strong track record of open-source scientific software development. PICSL was one of the founders of the National Library of Medicine Insight Toolkit (ITK) and is a major contributor to the most recent ITK version 4. PICSL leads the development of other significant open-source tools, including interactive image segmentation tool ITK-SNAP, deformable image registration package ANTS/SyN, and diffusion MRI analysis package DTI-TK. All three of these open-source applications tools are used widely by researchers, and are estimated to have thousands of users. This project will take full advantage of the software development infrastructure and expertise at PICSL. These best practices include hosting source code, documentation, and Wikis on public repositories such as GitHub and SourceForge. Concurrent development by multiple programmers is facilitated by the use of Git software. Cross-platform nightly builds are performed on a dedicated server that contains multiple VMware Fusion virtual machines running different versions of Windows, MacOS, and Linux operating systems. These virtual machines are automatically activated at night to compile the latest source code in the repository, perform automated testing, and upload binary executables to a public file download area. The results of the compilation and testing are sent to a web-based
dashboard. This build process relies on open-source CMake, CTest, CPack and CDash tools from KitWare, Inc.

**Office Resources**

PICSL currently occupies 2,000 ft² of space on Penn campus, and in the summer 2015 will move to a larger 2,400 ft² space in the renovated Richards building. Each investigator will have office space in this building, along with shared office and open space for trainees and students. Michele Haines, the dedicated administrative assistant for PICSL, will provide administrative support for the project.

**Resources for Medial Temporal Lobe Mapping Research**

**Penn Memory Center (PMC) and Penn Fronto temporal Dementia Centers (PFDC) Cohorts**

Building the combined *in vivo / ex vivo* MTL atlas in Aim 1 involves retrospectively accessing and analyzing *in vivo* MRI data from the records of patients evaluated at the Penn Memory Center (Dr. Wolk is the Assistant Director) and participants in imaging research studies conducted by both the Penn Memory Center and the and Penn Fronto temporal Dementia (FTD) Center, which is directed by Dr. Grossman.

The Penn Memory Center (PMC) is a single, unified Penn Medicine source for those age 65 and older seeking evaluation, diagnosis, treatment, information, and research opportunities related to symptoms of progressive memory loss, and accompanying changes in thinking, communication and personality. The PMC is a National Institute on Aging-designated Alzheimer's Disease Center (ADC). It is one of only 30 such sites in the nation. ADC designation is earned by leading universities and medical institutions offering state-of-the-science diagnosis, treatment, research, and more for individuals with Alzheimer’s disease, mild cognitive impairment (MCI) and other age-related progressive memory disorders. Research is a chief mission of the PMC. The PMC is located in the 2 South wing of the UPHS' new Perelman Center for Advanced Medicine (PCAM). The PMC is a multidisciplinary clinical and clinical research center providing state-of-the-art diagnosis and care for adults with cognitive disorders. It is designed for efficient research and clinical patient flow, comfort, and safety. The PMC currently includes six faculty clinician-researchers from 3 departments (psychiatry, neurology, medicine), two fellows, one resident, a nurse and other staff who provide patient care and conduct research studies. The PMC performs cutting edge patient-oriented clinical research. Currently, the PMC is conducting five NIH funded studies, 4 pharmaceutical industry clinical trials, and two private foundation studies of cognition in normal adults, mild cognitive impairment, Alzheimer’s disease and vascular cognitive impairment.

The Penn FTD Center is a state-of-the-art research center that brings together leading experts in neuropsychology, neuroimaging, clinical care, biomarker and cognitive neuroscience in an effort to improve the diagnosis, treatment, and care for individuals with FTD. In 2012, the Center evaluated over 500 patients. Currently, 240 patients participate in research projects that involve multimodal clinical, imaging, biofluid biomarkers, and extracted DNA data. The Penn FTD Center works in close collaboration with the Center for
Neurodegenerative Disease (CNDR) Neuropathology and Genetics Cores and with other neurodegenerative disease centers at the University of Pennsylvania. These include the Penn Memory Center, the Amyotrophic Lateral Sclerosis (ALS) Center, and the Morris K. Udall Center of Excellence for Parkinson’s Disease Research. Both centers operate multiple research neuroimaging studies, and the oblique coronal T2-weighted scan described in the Research Strategy is routinely acquired since 2010. The sequence was also incorporated into the clinical MRI protocol at the Penn Memory Center in 2011. As of January 2015, the Penn FTD Center had collected such scans in over 500 subjects, and the Penn Memory Center had collected them in over 800 subjects. Approximately 34% of the subjects participating in the T2-weighted scans consent to donate their brain to autopsy (50% of subjects at the Penn FTD Center and 60% of subjects involved in the National Alzheimer’s Coordinating Center (NACC) study at the Penn Memory Center, who represent ~35% of all scans at the Penn Memory Center). Autopsies are performed by CNDR pathologists and coordinated by the CNDR Brain Bank.

Center for Neurodegenerative Disease Research (CNDR) Brain and Biospecimen Banks

Tissue specimens (n=30) for the in vivo / ex vivo atlas proposed in Aim 1 will be obtained through a collaboration with the Center for Neurodegenerative Disease Research (CNDR). The CNDR brings together researchers investigating the causes and mechanisms of neurodegenerative diseases that occur more frequently with advancing age, including AD, Parkinson’s disease and other Lewy body diseases, frontotemporal degeneration diseases, amyotrophic lateral sclerosis and other motor neuron diseases. Its core and affiliated research programs encompass basic cell and molecular biology of protein misfolding and neurodegeneration, neurogenetics, neuropathology, biomarker discovery and validation, translational medicine and drug discovery, clinical research programs including the Penn Memory Center / Alzheimer’s Disease Center, and a database to integrate data from all of these programs on a common platform.

The CNDR established a biosample collection in 1989 to procure brains, cerebrospinal fluid, DNA, and plasma and serum from normal aged controls and patient subjects with neurodegenerative disorders. All specimens are tracked and linked with demographic and clinical data in a database and all samples and data are coded and de-identified to maintain patient confidentiality in accordance with HIPPA standards. Organ/tissue processing is optimized for multiple uses including confidentiality in accordance with HIPPA standards. All brain autopsies and diagnostic neuropathological examinations are performed in-house, under the direction of Dr. Trojanowski. Fixed and frozen tissues are currently available for research studies from more than 1,400 individuals. In addition, cerebrospinal fluid is available from >850 subjects and DNA/plasma/serum from >1,400

Small Animal Imaging Facility (SAIF)

Ex vivo MRI studies in Aim 1 will be carried out at the SAIF, which provides multi-modality radiological imaging and image analysis for cells, tissues, and small animals. The SAIF combines state-of-the-art instrumentation and a nationally recognized staff to assist
investigators with a wide range of imaging based experimental approaches. The SAIF currently provides a comprehensive suite of imaging modalities including: magnetic resonance imaging (MRI) and spectroscopy (MRS), optical imaging (including bioluminescence, fluorescence, and near-infrared imaging), computed tomography (CT), positron emission tomography (PET), single photon emission computed tomography (SPECT), and ultrasound (US). In addition, dedicated housing is available for mice and rats undergoing longitudinal imaging studies. Ancillary facilities and resources of the SAIF are devoted to chemistry, radiochemistry, image analysis and animal tumor models.

Ex vivo MRI will utilize a 9.4 Tesla (Varian, Inc., Palo Alto CA) 31 cm horizontal bore MR system (B100 John Morgan Building) equipped with a 21 cm ID 25 gauss/cm and a 12 cm ID 40 gauss/cm gradient tube and interfaced to an Agilent DirectDrive console. A custom M2M quadrature transmit/receive body coil with inner diameter of 35mm and a long z field (supporting B1 field of 80 mm FOV along the length of the specimen) was purchased in 2010 specifically to support human MTL tissue imaging. A number of other coils are also available.

Tissue preparation for imaging studies in Aim 1 will take place in a fully equipped animal surgery room adjacent to the Varian 9.4 Tesla MRI scanner in 101A John Morgan building. The scanner room also provides dedicated storage space our MTL tissue specimens.

**CHOP Pathology Core**

The scanning of histology slides in this project will be carried out at the Pathology Core Laboratory at the Joseph Stokes Jr. Research Institute in the Children’s Hospital of Philadelphia. In addition to a full range of histopathology services, the Core provides sophisticated imaging instrumentation for high-resolution whole slide scanning and virtual microscopy (ScanScope from Aperio). The PI has already collaborated with the CHOP core in the Phase 1 of the project. In the current proposal, the histology preparation and staining work will be carried out by through the subcontract to the Insautsi group at UCLM, whose techniques are highly optimized for anatomical and pathological imaging of the MTL. However, the ScanScope at the CHOP Core provides the best image quality and resolution, and will continue to be used in the proposed project.

**MRI Resources**

As of the summer 2015, the MRI facilities at Penn will include two research-dedicated 3 Tesla Siemens Prisma scanners and a Siemens 7 Tesla whole-body research MRI system. These scanners are operated by the Center for Advanced Magnetic Resonance Imaging and Spectroscopy (CAMRIS), which is responsible for maintenance, safety and staffing of human research MRI systems in the Department of Radiology. Maintenance includes service, upgrades and cryogen refills. A local Siemens engineer is available to address malfunctions. A CAMRIS committee reviews applications for MRI research studies on the Siemens systems for safety and feasibility and provides approval. CAMRIS collects an hourly usage fee, currently $500/hr, to cover the cost of scanner use, including maintenance and staffing by a certified MRI technologist. Additional support for MRI
Proposed MRI imaging for the aging study in Aim 2 will be carried out on a Siemens 7 Tesla whole-body research MRI system with a 32-channel head coil located in the basement of the Stellar Chance Building on the School of Medicine campus, a minutes walk from the Richards Building. This system also includes RF coils for 31P NMR spectroscopy and a custom hybrid head/neck coil for arterial spin labeling. A parallel transmission system is also available and can be used in conjunction with an 8-channel transmit/receive array. An outboard GPU processor has been interfaced with the 3T and 7T systems to allow rapid image reconstruction for multiband EPI and other high-throughput imaging.

Amyloid PET scanning with the F18 Florbetapir tracer in Aim 2 will take place at the Positron Emission Tomography (PET) Center at the University of Pennsylvania, one of the first PET Centers in the country. It was established in 1975 as a research facility for the study of human brain function. It continues to be a leading biomedical research institution with a large program in basic and clinical PET research. More than 5,000 PET studies are performed by the PET Center each year.

The PET Imaging Facility has four whole body PET/CT scanners, one Philips Gemini TF in the Hospital of the University of Pennsylvania and Philips Gemini TF Ingenuity, Philips Gemini TF BB, and Siemens mCT scanners in the adjacent outpatient building. Separate rooms are available for patient preparation and injection of radiopharmaceuticals. In addition, there are nearby hot labs for preparing and extracting doses and a blood lab for sampling and counting. The Philips Gemini TF Ingenuity will be used for this study. It has an open design that allows patients to see outside the scanner, which improves patient acceptance. It has an 18-cm axial FOV and 70-cm diameter gantry aperture. The intrinsic spatial resolution is 4.8 mm, with energy resolution of 12% for good scatter rejection, and very good system timing resolution of 500-550 ps, which enables time-of-flight (TOF) measurement for improved image quality and data quantitation. The TOF information along with physical data corrections (e.g. scatter, attenuation) are included in the system model of the iterative ML-EM list-mode algorithm which is used for image reconstruction. Attenuation correction and anatomical registration are provided by the CT scanner. The Philips Gemini TF Ingenuity is shared 50/50 between research and clinical examinations. Daily quality control is performed and system performance is maintained by the Physics Instrumentation Group in the Nuclear Medicine division.

A cyclotron and radiochemistry facility provides space for the JSW, BC 33015 cyclotron, and its accessories including targets for production of short-lived radionuclides as well as a radiochemistry laboratory for chemical syntheses by which product radionuclides are incorporated into trace molecules. The Facility includes: 1) Cyclotron vault, 2) Cyclotron maintenance workshop, 3) Control room, 4) Radiochemistry laboratory, 5) Mechanical room, 6) Power supply room, 7) Counting room.
Equipment

N/A

Specific research plan

Specific aims

We have entered an age of unprecedented scale, resolution, and accessibility of digitized biomedical image data. The BigBrain (Amunts et al. 2013) project is a publicly accessible digitized image data set of an entire human brain at a resolution approaching individual cells, painstakingly acquired and processed over the course of a decade, with a corresponding magnetic resonance image (MRI). The BigBrain offers an important opportunity to provide common data and standards to bring together neuroscience communities focusing on the macroscopic as well as the microscopic structure of the brain. Since the microstructural (cytoarchitectonic) boundaries cannot consistently be predicted from macroscopic boundaries (Amunts et al. 2005), the BigBrain should be anatomically labeled at multiple scales to maximize its potential. Unlike MRI data, manually labeling a single such brain at high resolution would take an expert many years to complete, and it takes years to train an expert on a given labeling protocol. With the prospect of more high-resolution biomedical images becoming available, we need to scale up our ability to label cytoarchitectonic data such as in the BigBrain. Facing a similar challenge, EyeWire is a successful computer game created for humans to segment mouse brain scanning electron micrographs into three-dimensional neurons. The primary objective of our project is to create a computer game for microstructural labeling of human brain imaging data.

Our proposal joins the expertise of the following people: Dr. Arno Klein (Sage Bionetworks) has expertise in brain image processing and has led manual and automated human brain image labeling projects; Dr. Alan Evans (McGill University) helped to create the BigBrain and leads a team interested in making it more useful and more accessible, Dr. Paul Yushkevich (University of Pennsylvania) is a leader in hippocampal labeling of histology and MRI data; Dr. Jesse Himmelstein (Centre de Recherches Interdisciplinaires) creates video games for research and education and makes online platforms for sharing games; and Amy Robinson directs the EyeWire project.

Aim 1: To evaluate non-expert labels, experts will label a small subset of BigBrain hippocampal data. To provide training and to evaluate results of a crowdsourced task, it is necessary to establish gold standards. We will have experts manually label a portion of BigBrain hippocampal regions to establish these gold standards.

Aim 2: To scale up labeling efforts, we will break the labeling problem into smaller tasks within a computer game. Just as the EyeWire game breaks data into small cubes within which isolated tasks are performed by non-experts, we will break BigBrain data into smaller pieces to make labeling a parallel, distributed effort. To engage non-expert labelers, we will map individual tasks to elements of a game that a human would enjoy.
playing. If this game fails to engage sufficient numbers of players, we will enlist players through Amazon’s Mechanical Turk.

Aim 3: Aggregate the results of game play to label the BigBrain. We will combine the thousands of results of playing the game to establish labels for the BigBrain hippocampal regions.

Exploratory Aim 1: Assess the potential for extending the game to all brain regions. The hippocampus is an important region to label, but ultimately we want to label the entire brain, so we will explore how our approach generalizes to all other brain regions.

Exploratory Aim 2: Train a supervised learning algorithm on the crowdsourced results. We propose to radically scale up our ability to label imaging data by training a supervised learning algorithm on the crowdsourced labels, and evaluate how well we can automate labeling.

Research strategy

(A) SIGNIFICANCE

The importance of segmenting the BigBrain:

The BigBrain (Amunts et al. 2013) represents a decade of work to construct a high-resolution 3-D model of a human brain at nearly cellular resolution (20μm isotropic) based on reconstruction of 7,404 histological sections stained for cell bodies (Fig. 1). In 2014, we extended our processing of the BigBrain to include improved section-to-section intensity inhomogeneity correction, 3-D tissue classification, and cortical surface extraction (Lewis et al. 2014). The BigBrain is extremely important to the neuroscience community because it represents whole-brain histological data that match its accompanying magnetic resonance imaging (MRI) data. It is the best brain imaging data set in the world to establish a canonical space at both microscopic and macroscopic resolutions. However, for the cell-stained histology data to be truly useful, it needs to be segmented into cytoarchitectonic regions, as has been the standard since Korbinian Brodmann (Brodmann 1909). Cytoarchitectonic labeling is important for neuroscience and human health:

- Delineation of regions on the BigBrain, whether obtained from a manual or automated approach, will help improve statistics in probabilistic maps of neocortical and subcortical areas, as well as fiber tracts (although the stain used was for cell bodies, some fibers can be followed in areas of high contrast).

- These cytoarchitectonic maps can be warped to a common reference frame and used as a topographical reference for anatomical localisation of activations observed in functional imaging studies, or as a high-resolution 3-D neuroanatomical guide (atlas) for microsurgery.
Labeling the cytoarchitecture of the human brain requires great expertise acquired over many years of labeling. It is a slow and tedious process; it would take an expert many years to draw all of the major cytoarchitectonic boundaries for the 7,404 sections of the BigBrain, and unless one could convince many such experts to do it, the necessary inter-rater reliability statistics of consistency cannot be measured. In other words, it will be a long time before we see the cytoarchitectonic boundaries of the BigBrain comprehensively labeled by experts, diminishing the tremendous potential that this data set has to offer. In this project, we propose to overcome the daunting challenge of labeling BigBrain cytoarchitectonic regions through crowdsourcing.

Labeling the hippocampus:

The medial temporal lobe (MTL) is a complex brain region of enormous interest in research on memory, aging, psychiatric disorders, and neurodegenerative diseases. Within the MTL, the subfields of the hippocampus (cornu Ammonis fields CA1-CA4, dentate gyrus, subiculum) and the adjacent cortical subregions of the parahippocampal gyrus (entorhinal cortex, perirhinal cortex, and parahippocampal cortex) are understood to subserve different functions in the memory system (Moscovitch et al. 2006, Bakker et al. 2008, Wolk et al. 2011). Different psychiatric and neurological disorders are known to affect MTL subregions differentially, selectively, and in a complex progression (Small et al. 2011, Braak and Braak 1995, Arnold et al. 1995, De Lanerolle et al. 2003, West et al. 2004, Lucassen et al. 2006). The non-uniformity of MTL involvement in normal brain function and in disease makes in vivo interrogation of the structural and functional properties of MTL subregions highly desirable. More studies are adopting high-resolution MTL imaging, creating a growing need for accurate and reliable computational techniques for analyzing these data.

Dr. Yushkevich (University of Pennsylvania) is a world expert in hippocampal segmentation and works with hippocampal image data at both scales reflecting BigBrain modalities -

---

Figure 1. From Amunts et al. (2013)
microstructure and macrostructure (see Fig. 2). His group has been a major contributor to
the MTL subregional imaging field, in particular to the effort to bring the MTL imaging and
neuroanatomical communities together with the goal of developing a harmonized protocol
for MTL subregion segmentation, and took a lead role on the first paper to emerge from
this effort (Yushkevich et al. 2015). In the proposed project, the UPenn team will provide
reference segmentation of hippocampal subregions in a subset of slices from the BigBrain
data set as gold standard labels for evaluating crowdsourced labels, toward a rich ex vivo
atlas that would greatly facilitate comparisons between different in vivo and ex vivo studies
of the hippocampus.

Figure 2.

Example hippocampal subfield labels.

Left: Example of hippocampal subfield labeling. Right: The first work demonstrating
hippocampal subfield labels in MRI space that are derived from ground-truth histological
imaging (Adler et al. 2014).

Computer games for solving scientific problems:

There has been increasing interest in and attention given to “citizen science” initiatives to
crowdsource the solving of scientific problems through what is sometimes termed “human
computation.” Compared to computers, humans currently excel at face and pattern
recognition, object identification, abstract symbol processing, etc. Examples of recent
scientific computer games which have attracted many players include Foldit and EyeWire
(Fig. 3).
Figure 3.

Screenshots of popular scientific games.

FoldIt, Nanocrafter, and EteRNA tutorials are important for training and evaluating players and for providing scientific context, so that the player is further motivated to play. EyeWire has nearly 200,000 players who build neural connections from mouse scanning electron micrographs.

a: FoldIt tutorial
b: Nanocrafter tutorial
c: EteRNA tutorial
d: EyeWire tutorial
e: Eyewire neural reconstruction with scoring leaderboard
**Foldit** is an online puzzle video game where the objective is to fold the structure of selected proteins using various tools provided within the game. Foldit’s players (now estimated at 240,000 people (Marshall 2012): provided useful results that matched or outperformed algorithmically computed solutions in 2010 (Cooper et al. 2010), helped to decipher the crystal structure of an AIDS-causing monkey virus in 2011 (Khatib et al. 2011), and achieved the first crowdsourced redesign of a protein in 2012 (Eiben et al. 2012). Some of the makers of Foldit have introduced two new scientific discovery games, one called **EteRNA**, in which a player designs RNA molecules to conform to desired shapes, and the other called **Nanocrafter** about synthetic biology, where a player uses pieces of DNA to build everything from computer circuits to nanoscale machines.

**EyeWire** is a game to help map neural connections in the visual cortex of a mouse brain played by nearly 200,000 people worldwide. The player does this by linking together computer-rendered, 3-dimensional segments of tissue extracted from scanning electron microscope sections to make neural processes that start from one face of a small cube of data to another face of the cube. By breaking the problem down from something currently computationally intractable (segmentation of interconnected neurons from SEM data) to one that is not only tractable but engaging for players (building a scaffold to escape a cube), they are able to crowdsource a very difficult problem. *Likewise, the primary objective of our project is to crowdsourcethe segmentation of brain image data, in our case the cell-stained histological sections from a human rather than SEM sections from a mouse.*

**HTML5 technologies for browser-based scientific games:**

The Web browser is doubtlessly the most widespread and cost-effective way to reach large numbers of potential players of scientific games. And yet, only a few years ago, technical difficulties prevented developers from delivering interactive and computationally intensive experiences over the Web. With the increasing uptake of the HTML5 standard by popular Web browsers, Web applications can now include 3-D graphics (WebGL), high-fidelity sound, and real-time communication with servers, all without browser plugins that require installation. Even better, many of these standards are being adopted by mobile Web browsers, meaning that the same application is available directly on a player’s smartphone or tablet.

The Centre de Recherches Interdisciplinaires (CRI) developed two technologies specifically created for scientists to take advantage of HTML5 technologies for games. **RedWire** (Fig. 4) is a new kind of game engine for remixing games. Game creators can take apart existing game designs, mix them with others, and add their own functionality. Scientists can create new games faster, and others can contribute variants of their games that may even outperform the original. The novelty of the engine lies in allowing developers to create imperative blocks of code within a functional framework, execute these blocks in parallel and merge the resulting changes, representing input and output as data buffers, and providing visual programming and debugging tools.
RedMetrics (Fig. 4) is the first open data, open source game analytics service. Once a game is released, its creators want to track how players interact with it. Knowing where players get stuck or give up is invaluable for improving the player experience and encouraging players to continue. For a scientific video game, it is critical that teachers and researchers can track how players interact with the game in order to follow the learning curve or gather experimental evidence. Since no open source solution currently exists to gather in-depth metrics on player behavior, RedMetrics is designed to fill that gap. As a Web service, it is capable of receiving information about any game, regardless of the platform it is developed on.

(B) INNOVATION

Aim 1, while focusing on traditional neuroanatomical methods for segmenting brain cytoarchitecture, is nonetheless innovative since it will modify a hippocampal labeling protocol for use with ex vivo hippocampal data, and will make use of these data as gold-standard labels for evaluation and scoring of players of a computer game.

Aim 2 is innovative since it will be the first instance of crowdsourcing cytoarchitectonic processing or analysis, and that too by individuals who have no neuroanatomical expertise (in contrast with Aim 1). Moreover, this crowdsourcing will take the form of a computer game, and will be the first to deal with cytoarchitectonic data. Players’ repeated segmentations will also be used to generate intra- and inter-rater reliability statistics for cytoarchitectonic segmentation, which will complement existing data on inter-brain anatomical variation of cytoarchitectonic boundaries (Amunts et al. 2005). The game will use the first open data, open source game analytics service (RedMetrics).

Aim 3 will aggregate results from the crowdsourcing above, and will constitute the first time cytoarchitectonic boundary delineation will have been undertaken on small portions of the
image for which (macrostructural) anatomical context has been removed. Another innovative contribution will be the software written to aggregate and reconcile tens of thousands of drawn curves to construct 3-dimensional boundary surfaces.

**Exploratory Aim 1** is innovative in that it proposes to generalize a game-based approach tailored to a specific domain (the hippocampus) to a larger context (the entire brain).

**Exploratory Aim 2** proposes the use of crowdsourced segmentations to train a machine learning algorithm to perform the same task. If this algorithm performs well, this will introduce a new and powerful approach to inferring cytoarchitectonic boundaries of brain regions.

**(C) APPROACH**

**Aim 1: To evaluate non-expert labels, experts will label a small subset of BigBrain hippocampal data.**

Dr. Yushkevich will oversee the manual segmentation of BigBrain hippocampal sections for use as a gold standard to evaluate each player’s accuracy and consistency, and will use this information to weight player contributions accordingly. Manual segmentation of the entire hippocampus at the resolution of the BigBrain is beyond the scope of this proposal. The UPenn team will label the left and right hippocampal regions in a subset of 60 slices each, taken in the coronal plane with approximately 1mm intervals. This represents 2% of the total number of slices spanning the hippocampus. We will store and release these segmentations and their boundaries as a publicly accessible project in Sage Bionetworks' Synapse platform.

*Segmentation Protocol :*

The segmentation protocols by Ding and Van Hoesen (Ding and Van Hoesen 2015) and Duvernoy (Duvernoy 1999) will be used as references for the BigBrain segmentation protocol. This protocol uses a combination of different histological stains (e.g., Nissl anatomical stain, AT8 tau stain, etc.) to define rules for distinct anatomical subfields in the hippocampus, with particular focus on the anterior hippocampus where the structure is very complex in shape and organization. Since the BigBrain only uses a single stain, the protocol will be modified to use the features available in the BigBrain data. Additionally, the UPenn group will stay abreast of the active effort to harmonize the hippocampal subfield segmentation protocols (Yushkevich et al. 2015). Laura Wisse, a postdoctoral fellow in the group is involved in the effort to develop harmonized boundary definition rules for the harmonized protocol. Although this protocol targets MRI, the boundary working group is also using histology data to test the veracity of proposed rules. A team of neuroanatomists is examining rules from prior histology and MRI segmentation protocols, and working to define rules that are consistent between them. When the rules adopted from Ding deviate from the rules put forward by this group of anatomists, the UPenn team will use the consensus rules to ensure better alignment with the harmonized protocol for MRI segmentation (expected in 2016-2017).
Boundaries:

Boundaries can be defined either by a difference in the size/shape/density of cells, or in the presence/absence of layers. The hippocampus is considered more primitive cortex and doesn’t have the distinct laminar appearance of the cortex. However, parts of the head of the hippocampus do appear this way, and this is one of the ways an expert uses to differentiate regions. The head of the hippocampus also has multiple classes of cells within each subfield. For example, CA3 cells appear quite different in the uncal, vertical, and typical regions (Ding and Van Hoesen 2015).

Substructures:

The segmentation protocol will label the following hippocampal subfields: Cornu Ammonis (CA) fields CA1, CA2, and CA3; Dentate gyrus hilar region (DG:H), which is also referred to as CA4 by some authors, and the dentate gyrus proper (DG); subiculum (SUB). The hippocampal layers stratum radiatum and stratum lacunoso-moleculare will be assigned a separate label (SRLM), which correspond to the prominent hypointense band in MRI images of the hippocampus.

Operationalization:

Segmentation will be performed by John Pluta, who has over 5 years of expertise in hippocampal segmentation including ex vivo MRI and histology. He will use the HistoloZee tool (see Fig. 2), developed specifically for working with large 3D histology stacks and their segmentation. He will perform segmentation on the highest available resolution data from BigBrain and will use histology data from UPenn and published sources (Ding and Van Hoesen 2015, Duvernoy 1999) for reference. Segmentations will be exported in vector graphics format allowing for future modification.

Reliability:

Inter-rater and intra-rater reliability for the protocol will be established. A set of 10 slices will be used for this purpose. John Pluta will resegment these slices after 1 month or more delay to establish intra-rater reliability. Laura Wisse will perform separate segmentations 1 month apart to establish her own intra-rater reliability and inter-rater reliability with Pluta. Reliability will be reported using the intraclass correlation coefficient, Dice similarity coefficient (a relative overlap measure) and boundary distance measures. The intra- and inter-rater reliability will be valuable data for answering hypotheses about the ability of non-expert gamers to replicate the work of anatomists.

Aim 2: To scale up labeling efforts, we will break the labeling problem into smaller tasks within a computer game.

We will break BigBrain data into small pieces to turn labeling into a parallel, distributed effort. Labeling each piece would constitute an element of a game played by non-experts.
Seed boundaries:

The expertly drawn hippocampal subfield label boundaries will only cover a small portion (2%) of the BigBrain’s hippocampus (and none of the rest of the BigBrain). We can supplement these with less accurate, automated label boundaries to get a rough sense where we should expect these boundaries to lie, by affine registering other labeled image volumes to a downsampled version of the BigBrain. We anticipate that the inter-subject anatomical variability (Amunts et al. 2005) in the different images and the affine transformation will result in a set of non-overlapping “seed” boundaries in the space of the BigBrain. We can use these seed boundaries to select image tiles and to provide hints while playing the game.

Image tiles:

We will first present the labeling problem to players much as it is presented to expert labelers of neuroanatomy: as 2-D images upon which cytoarchitectonic boundaries are to be drawn (Fig. 5). Beyond this, the experience will be quite different. We will not present an entire section to the player, but instead present a small portion (tile) containing gray matter, so there is no context of where in the hippocampus it came from. We will only show players image tiles that intersect seed boundaries, and for each original tile, we will generate two additional tiles translated in x or in y by half of the tile’s width. This sliding window approach to tiling the image will account for cases where a label boundary lies along a tile’s edge. Generating the tiles and serving them to remote clients will be done using the current BigBrain tile server mechanism developed by Dr. Evans’ team. A multiresolution pyramidal 3D structure is generated with blocks of a chosen size from which remote clients can obtain tiles from arbitrary spatial plans at the resolution matching the proper zoom level. The gaming application will only require a subset of this flexible data structure; we will choose the appropriate spatial plans, tile size and resolution to present to the gamers. The current system and infrastructure, hosted on the CBRAIN platform (Sherif et al. 2014), has been benchmarked at 16Gbps for extremely rapid tile read operations during real-time visualization, which is sufficient to serve single images to many concurrent gamers.

Scoring players:

Players would draw a line or curve on an image tile, and would be scored in the following ways:

1. **Accuracy**: similarity to an expert’s drawing
2. **Intra-rater reliability**: similarity to previous drawings by the player on the same tile
3. **Inter-rater reliability**: similarity to other players’ drawings on the same tile
4. **Registration-based reliability**: similarity to the seed boundaries used to select the tile
5. **Independent reward**: a score that is independent of other lines
For 1-4 above, similarity between two drawings can be computed by the Hausdorff distance measure, or similarity between regions delineated by the drawings can be measured by various overlap measures (Klein et al. 2009). An independent reward can either reflect drawing speed or some other aspect of game play that has nothing to do with the line, such as discovery of a random treasure. This could be used to motivate a player to contribute drawings from which intra- or inter-rater reliability measures can later be computed.

**Training:**

A new player will have no idea how to draw a boundary between cytoarchitectonic regions, and will rely on the game for initial training and assessment. A proven technique in these circumstances is to present players with “reference” puzzles where the solution is already known (see Fig. 3). In this case, the game will show a player image tiles that have been previously labeled with cytoarchitectonic boundaries by experts, although the player does not see the previous labels. The player will draw a boundary and will receive an accuracy score, and based on the score, may receive further training. During actual game play, every so often an image tile from an expertly labeled section will be presented to the player to evaluate how well the player is performing, and perhaps to infer effects of learning, attention, and drift. These intermittent evaluations can be used to retrain the player, or weight the confidence in their contributions. Since the game will compute intra-rater reliability, inter-rater reliability, and registration-based reliability for every drawing, these scores can be used as a proxy for accuracy, or to determine whether the player should be evaluated for accuracy.

**Game mechanics:**

There are a number of possibilities for the central game mechanic. One promising idea is to represent the labeling process as a drawing or painting game. This approach could...
appeal to the artistic side of players, especially if the game supports touch controls on smartphones and tablets. Another idea is to approach the labeling as a game about cutting or slicing, which is both a popular mechanic among mobile games and may appeal to players as being akin to brain surgery. The “surgery” would introduce a goal that is independent of labeling: the player would find a path to reach a tumor that must be removed. If the path deviates too much from the “correct” path, the player would be penalized, as in the popular “Operation” game currently manufactured by Hasbro, whereas if the path is close, the surgery is successful and another patient is presented to the player. A “correct” path could reflect one of four scores above: accuracy, intra-rater reliability, inter-rater reliability, or registration-based reliability.

With any (or all) of the types of scores, it would be possible to set up the game as a competition among players, and to display a leaderboard with players’ highest scores. Levels could be achieved based on labeling difficulty, determined by intra-rater and inter-rater reliability statistics obtained for expert labelers or for all players. Alternatively, the game could reward cooperation among players. For example, players could try to guess where others have drawn their lines on a given image tile. This would promote inter-rater reliability and therefore more consistent labels. FoldIt and EyeWire have demonstrated the power of social tools within citizen science games. In addition to a cooperative mechanic, we will consider bringing social interaction to our game via chat, team play, or badges.

The power of crowdsourcing lies not just in the number of minds focused on a challenge, but in the different and unexpected solutions that can arise. Rather than assume that the original image data will be conducive to drawing boundaries, if we were to make available browser-based image filters to enhance contrast, alter colormaps, etc., players could explore different ways of seeing and segmenting cytoarchitectonic regions. The Cancer Digital Slide Archive (CDSA) tool (below) for browser-based visualization of high-resolution image data such as the BigBrain has image filtering and semi-automated segmentation capabilities. We will explore different ways of enabling players to filter the image tiles and evaluate whether image filtering improves labeling accuracy.

Both the drawing and slicing mechanics above offer players the enjoyment of recognizing patterns and the sense of completion that comes with finishing a puzzle. These mechanics could be embellished by adding secondary mechanics such as time limits, treasures to discover, territory to surround and conquer, enemies to fight off, etc. For better or for worse, these “flashier” mechanics turn the focus away from the central scientific task, which may help to keep some players engaged and may turn others away who were attracted by the appeal of a game that purports to contribute to science and society. For this reason, we will need to strike a balance to discover which game mechanics best motivate players to contribute to the game.

Game development:

We will begin our game design process by prototyping different game mechanics (above) and testing them directly with players in order to evaluate which ideas provoke the most engagement and provide the best results. Building on this initial insight, we will use an agile
software development approach to develop a polished game that we can test with larger and larger user groups. This will resemble the iterative process of design adopted by the FoldIt developers (Cooper et al. 2010).

The ubiquity of the Web browser makes it the platform of choice for a citizen science game that wants to lower the barrier of entry to new users. We will build our game upon JavaScript and HTML5 technologies, ideally without additional plugins that require installation. This approach will enable us to draw upon Dr. David Gutman’s CDSA image viewer (example viewing BigBrain data: http://node15.cci.emory.edu/BIGBRAIN/), Montreal Neurological Institute collaborator Dr. Alan Evans’ BrainBrowser, Dr. Roberto Toro's Seadragon-based drawing tool, and/or the RedWire game engine, as needed.

Data handling:

Each image tile would have a unique identifier, as would each player, and each player’s drawing. This information would be stored in a CouchDB or PostgreSQL database management system. As players play the game, image tiles would be loaded into the game (as described above), and players’ vector-based drawings would be stored in the database as JSON objects. Regarding player interaction with the game, we will link the game to the RedMetrics open game analytics service, so that such data will be accessible to citizens, scientists, and game developers alike for analysis.

Aim 3: Aggregate the results of game play to label the BigBrain.

As described above, the proposed game will result in multiple manually drawn curves for each image tile, each one corresponding to a putative cytoarchitectonic boundary for a small portion of a given histological section containing hippocampus. Aggregation of the large collection of manually drawn curves will involve the following steps:

1. Position the multitude of curves in the original space of the BigBrain.
2. Weight the confidence in each curve based on the evaluated skill or estimated performance of each player at the time it was drawn.
3. Propagate label boundary assignments from expert-labeled curves to all other curves.
4. Create a 3-dimensional, probabilistic label assignment for each volume element (voxel) based on its position with respect to all label boundaries in 3-D.
5. Create an additional atlas with hard boundaries based on label fusion (Fig. 6, Wang and Yushkevich 2013), or on a modified version of the STAPLE algorithm (“Simultaneous Truth And Performance Level Estimation”) that attempts to estimate ground-truth labels based on incomplete labeling by multiple labelers (Landman et al. 2010).
Exploratory Aim 1: Assess the potential for extending the game to all brain regions.

Ultimately we want to label the entire brain, so we will explore how our approach generalizes to other brain regions. Since we have expert manual labels drawn by Dr. Evans' team for some limbic regions, we will start with these as gold standard data within the game. If there is time, we will evaluate how similar players are at delineating other regions for which we do not have expert labels. For example, we can test to see how consistently players assign boundaries for cortical regions throughout the BigBrain, and see how these boundaries compare with expected number and position of Brodmann label boundaries (Brodmann 1909).

Exploratory Aim 2: Train a supervised learning algorithm on the crowdsourced results. To explore the potential for learning from crowdsourced human results to automate labeling, we will train a supervised learning algorithm. We will use supervised machine learning methods in the open source scikit-learn Python package, the open source Dato Python package, and our own in the R programming environment. We have considerable experience building and training classifiers and regression models, and are very interested to try deep learning (convolutional neural network) approaches using the Dato or Theano Python packages. We have previously applied deep learning using Theano to automatically segment human brain MRI data at a macroscopic scale, and are in communication with developers of Dato for guidance in applying deep learning to the BigBrain's microscopic data.
Contingency plans and timeline:

Fig. 7 provides a timeline to achieve this proposal's goals.

![Timeline](image)

Figure 7.

**Timeline.**

We will prepare the gold standard evaluation data (Aim 1) and develop and update the game (Aim 2) through the 18th month as we get feedback on its use. Year 2 will consist primarily of testing the aggregation of segmented data (Aim 3), exploring how well the game can generalize to segmentation of every region of the BigBrain (Exploratory Aim 1), and training and testing an automated approach that learns from the crowdsourced data (Exploratory Aim 2), as well as to publish and present our findings.

Without training or initialization, it is clear in Fig. 5 that a player would simply draw a line between a darker and a lighter region, or between layers, rather than through gray matter for this difficult boundary. The orientation can be initialized in several ways to avoid these tendencies, such as by providing a “starting line” on the outer edge of the hippocampus or by providing a hint, such as the seed boundaries used to select the tile.

If players are unable to draw consistent boundaries, we will increase their training against gold-standard evaluation data and provide hints from better players and from co-registered label boundaries. If this doesn't work, we can simplify the task and ask players if there is, or is not, a boundary within a given tile. A game that can generate a consistent set of boundary-containing tiles will be a success, because we will be able to infer the probabilistic positions of label boundaries, which may be sufficient for most label-based applications, or could be used to help guide expert labelers so that they are much more efficient.

If we fail to attract many players, we will present the game through Amazon’s Mechanical Turk (AMT). AMT provides an online, on-demand, scalable workforce, where each “human intelligence task” (HIT) is submitted by a Requester and is performed by one of tens of thousands of paid Workers around the world. AMT has been used in many scientific experiments, including cognitive behavioral experiments (Mason and Suri 2012, CogSci2011 attendees 2011), and is considered a convenient, affordable way to attract many workers to perform tasks that a computer is currently poor at executing. The median wage is approximately one to two dollars per hour, and short tasks (around five minutes) are awarded around 10 cents. We would pay Workers a penny to play each round of the game. It would be straightforward to turn our game into AMT HITs using the open source psiTurk Python library (http://gureckislab.org/mtworkshop/), an open platform for setting up,
testing code, posting HITs, and paying Workers on Mechanical Turk. We would also share our open source code under an Apache v2.0 license with anyone via GitHub and through psiTurk’s Experiment Exchange. We have considerable experience designing and building Web apps for visualizing and interacting with data, and are poised to modify the browser-based game so that it is accessible to AMT Workers.

Project

The principal aim of this proposal is to crowdsource the segmentation of brain histological data, specifically the cytoarchitectonic regions of the hippocampi of the human brain, by means of a computer game. Currently, only human experts perform reliable cytoarchitectonic labeling at a very slow and small scale, whereas we propose to enlist many human non-experts to engage in a distributed version of this task at a quick and large scale. By turning this arduous, isolated task into an engaging, collective activity, we hope to radically change the way anatomists approach segmentation/labeling.

To support this aim, we must (1) prepare expert (gold standard) labels to a subset of the hippocampal sections to evaluate crowdsourced results, and (2) aggregate the crowdsourced results to label the hippocampi. For our first exploratory aim, will explore how our approach generalizes to all other brain regions, and for our second exploratory aim, we will train a supervised learning algorithm on the crowdsourced results and evaluate how closely the automated approach matches human assessments.

We will use the BigBrain, a high-resolution 3-D model of a human brain at nearly cellular resolution (20μm isotropic) based on reconstruction of 7,404 histological sections stained for cell bodies. The BigBrain is extremely important to the neuroscience community because it represents whole-brain histological data with accompanying magnetic resonance imaging data. It is the best brain imaging data set in the world to establish a canonical space at both microscopic and macroscopic resolutions. However, for the cell-stained microstructural data to be truly useful, it needs to be segmented into cytoarchitectonic regions, which makes it the perfect focus for this project. Labeling of the BigBrain will establish an ex vivo atlas, a common space for neuroimaging data whose labels will provide a consistent, convenient, and meaningful way to communicate, classify, and analyze biomedical research data set in that space.

Call

Big Data to Knowledge (BD2K) Advancing Biomedical Science Using Crowdsourcing and Interactive Digital Media (UH2)
Hosting institution

Sage Bionetworks

Ethics and security

This proposal makes use of publicly available data.

Author contributions

AK conceived of and wrote this proposal.

References


A game for crowdsourcing the segmentation of BigBrain data 31

