

A Comparison of Neuroimaging Software and a Contour Inference Method for analysis of Task-fMRI data

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Declarations

I, Alexander Bowring, hereby declare that except where specific reference is made to

the work of others, the contents of this dissertation are original and have not been

submitted in whole or in part for consideration for any other degree or qualification

in these, or any other Universities. This dissertation is the result of my own work and

includes nothing which is the outcome of work done in collaboration, except where

specifically indicated in the text.

• The work presented in Chapter 3 has been published in the Neurolmage jour-

nal (Bowring et al., 2018). This work was presented at the Organization for Hu-

man Brain Mapping (OHBM) Annual Meetings in 2017 and 2018. At the OHBM

2018 Annual Meeting, this work was the recipient of an oral presentation and

Merit Abstract Award.

• The work presented in Chapter 4 has been published in the Neurolmage journal

(Bowring et al., 2018). This work was presented at the OHBM Annual Meeting

in 2017, where it was the recipient of an oral presentation.

• The work presented in Chapter 5 is based on a pre-printed manuscript.

Alexander Bowring

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Abstract

Over the last three decades, Functional Magnetic Resonance Imaging (fMRI) has rapidly progressed to become the primary tool for human brain mapping. Recently however, considerable attention within the field has been directed towards data-sharing and open science initiatives. This has been driven by a growing apprehension about the reproducibility of findings within the neuroimaging literature, amid concerns that current inference procedures are often misused or misinterpreted such that the overall scientific conclusions become distorted. One aspect specific to neuroimaging pinpointed as a cause for poor reproducibility is the high flexibility of a typical fMRI workflow. In the first part of this thesis, we investigate how the choice of software package used to conduct a statistical analysis can influence the group-level results of a task-fMRI study. We use publicly shared data from three published taskfMRI studies, and reanalyze each study within the three main neuroimaging software packages, AFNI, FSL and SPM, using parameteric and nonparametric inference. All information on how to process, analyze, and model each dataset we obtain from the publications. We use a variety of quantitative and qualitative comparison methods to gauge the scale of variability in our results and assess fundamental differences between each software package. While qualitatively we find broad similarities between packages, we also discover marked differences, such as Dice similarity coefficient values ranging from 0.000 to 0.743 in comparisons of thresholded statistic maps between software. We discuss the challenges involved in our replication attempt, while also utilizing open science tools in an effort to make our own research reproducible. In the second part of this thesis, we extend a contour inference method initially proposed by Sommerfeld, Sain, and Schwartzman (2018) (SSS) to develop spatial confidence sets (CSs) on clusters found in thresholded blood-oxygen-level dependent (BOLD) effect size maps. While traditional inferences based on hypothesis testing indicate where the null, i.e. an effect size of zero, can be rejected, the CSs give statements about where effect sizes exceed a positive threshold analogous to confidence intervals simultaneously across the entire brain. We make advancements to theoretical aspects and implementation of contour inference to improve the method's finite-sample performance. We extend the wild bootstrap theory presented in SSS, proposing a method based on the t-bootstrap, and recommend that the bootstrapped residuals are multiplied by Rademacher variables instead of Gaussian variables. We also develop a linear interpolation method for computing the topological boundary over which the bootstrap is applied. Notably, we demonstrate that

the framework used in SSS for assessing simulations manifests considerable positive bias in the simulation results, and propose our own novel construction to solve this issue. In the final part of this thesis, we make further theoretical developments to contour inference so that the method can operate on the Cohen's d and partial R^2 effect sizes commonly reported at the end of a neuroimaging study. For the second and third parts of this thesis, we carry out intensive Monte Carlo simulations on synthetic 3D data to investigate the accuracy of contour inference on signals representative of fMRI activation clusters. We also demonstrate the method on two 'big' fMRI datasets, obtaining confidence sets to localize activation in functional data from the Human Connectome Project and UK Biobank.

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CHAPTER 6
Conclusion and Future Work

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