Modelling the effects of domestication in Wheat through novel computer vision techniques

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This report was submitted as partial fulfilment of a BSc degree in Computer Science (G401)

Declaration of originality

I confirm that:

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- I understand that there are severe penalties for Unacceptable Academic Practice, which can lead to loss of marks or even the withholding of a degree.
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Introduction, Analysis and Objectives

This project aims to answer a biological research question through the use of computer science, whilst also creating a software suite which will enable further studies to be carried out with ease.

Primarily the focus has been on the data science elements of my degree, creating, cleaning and discerning meaning in it.

Using a population of genetically diverse wheat, several hypothesis and questions are explored in the hopes of contributing to the scientific understanding of domestication. A mixture of image analysis through three-dimensional micro-computed tomography and computational analysis are used to provide these much needed solutions.

Additionally, as this is very much multi-disciplinary research, specific terms and definitions have been outlined in the *glossary* (table:7.3).

Background

Western society and agriculture has been dominated by the ability to create successful crops for the past 10,000 years [2]. Of these crops wheat is considered to be one of the most vital and is estimated to contribute to 20% of the total calories and proteins consumed worldwide, and accounts for roughly 53% of total harvested area (in China and Central Asia) [3].

During domestication, the main traits selected for breeding were most likely plant height and yield. This meant that important non-expressed traits such as disease resistance and drought tolerance were often neglected and lost overtime.

Whilst the choices made for selective breeding were successful, effects are now being felt as it is estimated that as much as a 5% dip is observed yearly on wheat production [3]. This decrease in efficiency is attributed to climate change bringing in more hostile conditions, which these elite and domesticated genotypes are unprepared for.

Furthermore, with increasing populations and less arable land there is an even greater pressure for the optimisation of grain and spike characteristics. With studies showing that spikelet count can be controlled by specific and sometimes recessive genes [4], which could drastically enhance overall yield, and a general public distrust towards genetically modification [5, 6, 7] the reliance on breeding programs for optimisation is further stressed.

Modern breeding programs have had some success in selecting primitive undomesticated genotypes and using them to breed back in useful alleles which would have been lost during domestication [8].

As such, there are questions still left open about how best to make selections for crop breeding. There is also a lack of formalised modelling of information which could be of use to these areas of research.

Biological Question and Materials

The driving question for this research asks "Can μ -CT data be used to model domestication in wheat?". Using an already grown and harvested range of genetically diverse wheat this project has generated a collection of 3D images, processed these images into raw phenotypic data and produced biologically significant information.

The genotypes used in this study are listed here, denoted by "X N" where X indicates the ploidy. 2N - Diploid; 4N - Tetraploid; 6N - Hexiploid.

• Wild Monococcum (2N)	• Durum (4N)	• Spelta(6N)
	• Dicoccoides (4N)	• Aestivum (6N)
• Domesticated Monococcum (2N)	• Dicoccum (4N)	• Compactum (6N)
(=- ')	• Ispahanicum (4N)	
• Tauschii (2N)	• Timopheevii (4N)	

Full species names are found in table:7.4.

Why use μ -CT image analysis?

In the past, science has been greatly limited by the amount of data which could be processed in an experiment. In the last few decades the inclusion of computer science has reduced this bottleneck. Now, the challenge for many fields of research is producing more data and this is often cited as the major bottleneck in creating robust studies [9].

Many experiments aim to meet the demand for data by using high-throughput automated imaging systems [10, 11, 12]. These systems have, in the last decade, become a standard and accepted tool for data generation. However, they will only produce 2-dimensional data on a per-plant basis. Image processing research has had success in modifying these automated systems in order to produce a pseudo 3-dimensional structure using stero-imaging [13]. Even so, these techniques require destructive harvesting of materials and do not provide information of internal structure.

For decades medical research has found success with X-Ray imaging technology [14]. From this, plant science has been able to benefit from the wealth of prior knowledge and more and more studies are being augmented with the use of X-Ray/ μ -CT imaging [15, 16, 17, 1, 18].

In this study, µ-CT has enabled the study of individual seeds of wheat, which is the product that plant breeders, commercial growers and farmers are truly interested in. Other imaging techniques could not provide as much detail, or in such a high throughput or quality.

Extracted Data

• Width

These samples come from over 70 plants and provided in excess of 2000 seeds for analysis which data was created based on. The traits recorded are labelled in figure:1.1 and are as follows:

auca	basea on.	The transferorded	are labelled in fig.	arc.i.i and arc	as ionows.	
• L	ength			• Volume		

• Surface Area

• Depth • Crease Depth / Volume

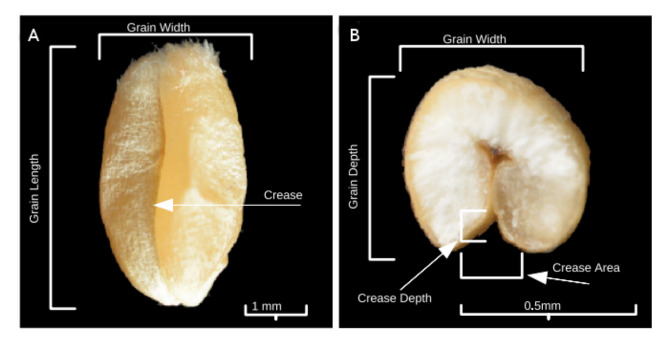


Figure 1.1: Wheat grain labelled (left), wheat grain cut in half (right), adapted from Hughes et al. [1]

Significance to Current Research

The biological interest in this area has been expressed in several areas of research [19], it is proposed that the key to unlocking diversity in the wheat genus lies in these ancestor, undomesticated species [20].

This research has the potential to be useful in several areas including: crop breeding; disease resistance; environmental stress. Each of these areas depend on making informed decisions in order to direct experiments. By producing information at an individual seed level, this study has been able to provide data that can offer suggestions of plant potential and behaviour.

Often, the most sought after traits are centred around thousand-grain-weight (TGW) as well as standard deviation of seed shapes. During harvesting, filters are used to only allow ideal shaped seeds through. This means that, potentially, despite a breed of wheat providing a high average volume of seed in reality much of it may go to waste if the shapes are not uniform. This research aims to alleviate this problem and provides low level information which is sorely required.

The individual images in figure:1.2 show, at a glance, the diversity and also the difference in the wild and cultivated (domesticated) species. This work allows for these differences to be quantified and evaluated into useful metrics for answering research based questions.

By better understanding the morphometric deviations in wheat species, more informed choices can be made when it comes to breeding wheat for the future and to fulfil ever-changing requirements.

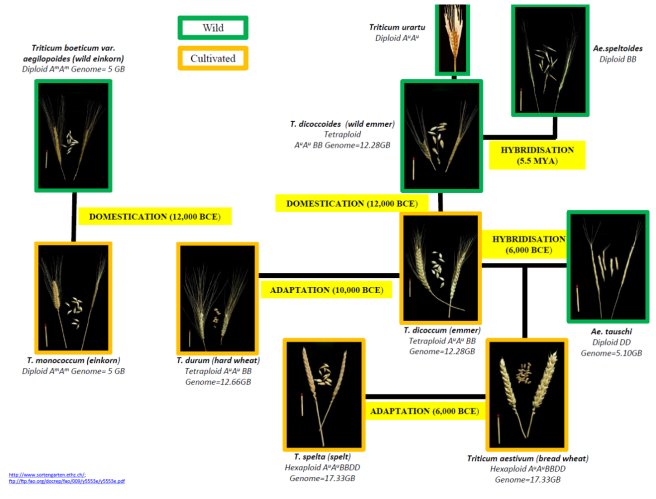


Figure 1.2: Phylogeny of wheat genotypes (Provided by Dr. Hugo Oliveira)

Aims and Objectives

The overarching aim of this project has been to create several pieces of software which aid in answering the biologically significant questions outlined. As well as to prove/disprove the hypothesis stated below.

The software created is robust in order to duplicate results and is flexible as to allow for further studies to be carried out and to use the same method.

Novel additions have been made to existing image analysis libraries in order to make them more flexible for this project. Figure:1.3 illustrates the range of diversity

Furthermore, the library written allows for easy data organisation and automation of otherwise difficult tasks such as concatenating data from multiple sources and graphing of information. Full documentation and integrated testing allows for a suite of tools which can be built upon in future and reduce the amount of effort required for similar studies to be carried out and analysed.

These aims have a focus on the phenotypic attributes generated from customised image analysis software [1] and can be seen in figure:1.1.

Hypothesis

To provide a full spectrum of analysis the null-hypothesis of this work is presented as investigating if there are morphometric differences in the seeds of several wheat varieties outlined in figure:1.2.

The comparison pairs are as follows:

- 1. Monococcum Wild and Monococcum Domesticate
- 2. Dicoccoides and Dicoccum
- 3. Spelta and Aestivum
- 4. Dicoccum and Durum
- 5. Monococcum Wild and Dicoccoides



Figure 1.3: Scans of wheat, showing diversity in Population, Compactum (6N) left, Durum right (4N)

Challenges Overview

The challenges which this project tackles come in two flavours: Computational and Biological. As such keen awareness of these is needed to appreciate the novelty of this work.

Biological Challenges

Previous studies have been able to demonstrate that variation in wheat grain morphology can be partially explained, in 2010 Gegas et al. demonstrated this through a 99.4% 2 component PCA [21]. However there is much left to do in terms of formal classifications and descriptions of these differences. This project deals with this problem through computational analysis.

Two effects run parallel in this study which requires acute biological knowledge of in order to make correct decisions:

- 1. The effects of ploidy in wheat.
- 2. The effects of domestication in wheat.

Hypothesis are required to take into account, both of these effects so as not to misidentify results.

Computational Challenges

Using µ-CT data in plant sciences is becoming more and more common [22, 15, 1, 17] and whilst a lot of studies focus on the traits of grains specifically no formal model has been created, no accepted data format. This is a data engineering problem and the methods described in this project address this.

Further to data organisation, proposals are made for the statistical analysis which should be used. This allows for studies to become more robust and repeatable, thus strengthening the studies overall.

The biological material used in this research is much more diverse a population than has been previously studied with μ -CT image analysis, this requires current computer vision methods to be adapted in order to be accurate.

Deliverables

This project provides three final deliveribles:

- 1. A flexible software suite written in Python that provides a standardised method for analysing and interpreting μ -CT data output.
- 2. A Graphical User Interface (GUI) which offers a point and click method for data gathering, graphing and manipulating μ-CT data, using the library from deliverable 1 as a backend.
- 3. Answers to the proposed questions (hypothesis), the *Results* and *Discussion* sections of this report provides this.

Software Design, Implementation and Testing

This chapter outlines choices and methodologies employed in the software engineering aspect of this project, as well as highlighting the key functional requirements and implementation decisions.

Functional Requirements

Requirements for this project are split between software requirements for both the CT Analysing Library and the CT GUI Application and the research requirements (i.e. the answers to the proposed hypothesis). Here the requirements for the software are discussed:

Requirements for CT Analysing Library

These are the functional requirements for the Python library produced:

- Provide an OOP means to deal with data
- Make gathering of data simplified
- Handle Saving of data in a useable format
- Easily enable data transformations
- Perform hypothesis testing

- Process rejoining of split scans
- Handle Removing of erroneous data
- Enable matching data to external information
- Auto plot data (boxplots, histograms etc.)
- Allow easy filtering of data

Requirements for CT GUI Application

- Provide a intuitive user interface for working with CT data
- Allow a interaction with data without the need for programming
- Implement the Matplotlib plotting utility
- Easily join experiment data with CT data
- Use an MVC model

- Implement the CT Analysis Library
- Display data visually
- Dynamically create graphs
- Provide hypothesis testing

Software Development Methodology

This project made use of formal design methods and strict organisation whilst being flexible to change. Overall the design took a hybridised form in order to best suit the scientific environment which this domain specific software is built for.

Data analysis drove the direction of the project, as a result an agile methodology was adopted. Weekly sprints were implemented as a list of "todo's", these were written on a Monday morning based off of the previous week's list.

Critical self-evaluation was performed by means of a "one-man SCRUM" meeting, this is a technique which requires self-discipline in order to accurately find faults and areas for improvement [23].

Further to this, regular meetings with research staff, at the National Plant Phenomics Centre, allowed for a developer-client relationship which SCRUM defines as being key. During these meetings details of the research was discussed and ideas given as to how future experiments could proceed. This allowed for critical decisions to be made as to software design and overall structure.

Language Choices

Both the CT Analysing Library and the CT Analysing GUI are implemented using the Python programming language, it has been developed and tested in versions 3.5 and 3.6 (Python 2 is not supported at all by this project).

In scientific programming three of the most commonly used languages are Python, R and MATLAB [24].

These three languages are able to provide all the features which this project requires. However Python was chosen for several reasons.

MATLAB could not be used as a potential language due to it being pay to use software, as this project aims to be accessible, the cost of software would greatly reduce the scope of access.

R is a valid candidate, it provides all of the statistical capabilities required by the project, it also provides packages for creating GUI based applications, it is fast and it is widely used in scientific computing and data science.

The main deciding factor is Python's wealth of resources, adoption rate and the developer of this project being vastly more experienced with Python's ecosystem than R's.

Designing Process

Through meetings and emails, the agile principles of communication over comprehensive documentation was used. Where conversations were decidedly much more beneficial than complex planing prior to developing a product.

Graphical elements, such as the graphing functionality of the CT Analysing Library and the CT GUI Application were sketched using wire-frames whilst in meetings where the potential users (clients) could provide their ideas.

In figure:2.1 an example of the wire-frames created during meetings is show (A), next to it is displayed the final look of the loading window (B).

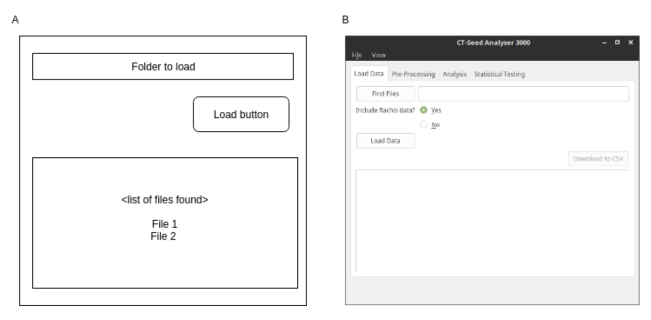


Figure 2.1: Wire-frame of the GUI loading data window

Similarly, figure:2.2 provides the initial wire-frame (A) of how the analysis window could have looked and what kind of GUI elements would be required, again, next to it is the final analysis window (B)

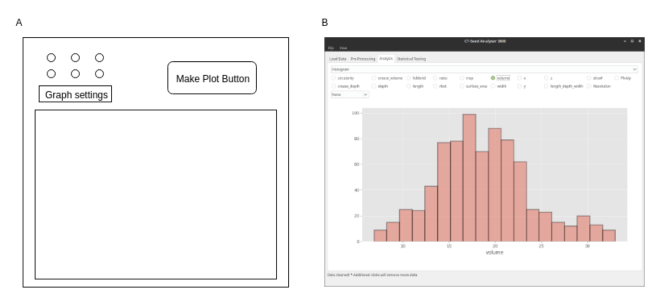


Figure 2.2: Wire-frame of the GUI analysis window

Documentation

Whilst an agile approach was used, some documentation was created for use with the CT Analysing Library.

The provided CT Analysing Library comes with "human-readable" format. Where most documentation generators (Doxygen, Pydocs, Javadocs etc.) implement very well structured and comprehensive documentation, the output is generally not very friendly and easy to read. Particularly for non-career-programmers. A

core feature of these provided software implementations are that they are well suited for a biologist, researcher or statistician to use.

This documentation generator was purpose created, implemented in LISP and provided in listing:2.

Beyond this, inline commenting is provided for supplied software. Keeping in line with the agile development ethos the software is self-documented and self-evident. A brief example of this is shown in listing:3

Documentation for the CT GUI application is provided as a visual user guide, and provides sample data for the user to test with.

Software Library Choices

Implementation

Design Patterns

CT Analysing Library Design Pattern

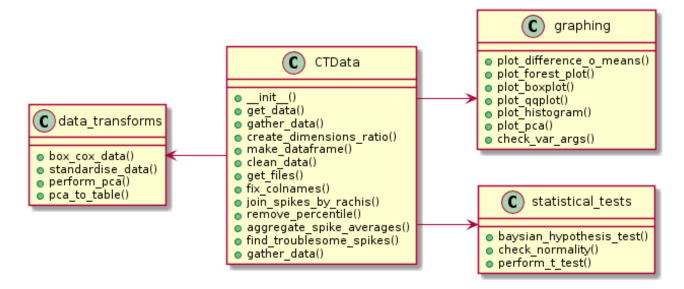


Figure 2.3: CT Analysing Library UML

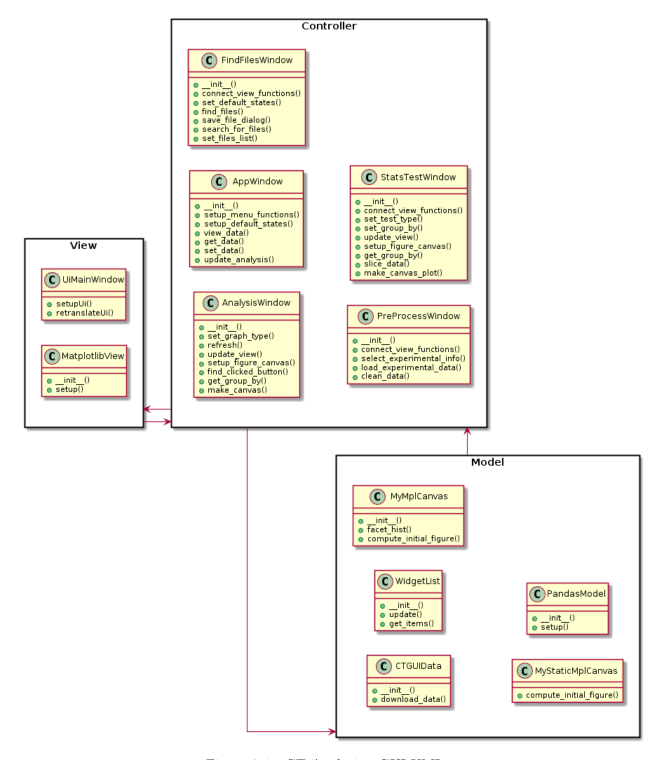


Figure 2.4: CT Analysing GUI UML

CT GUI Application Design Pattern

Version control

Testing

Feedback Forms

Feedback and constructive suggestions were made by researchers at the National Plant Phenomics Centre, these were submitted via the Google forms service. . .

Unit Testing CT Analysing Library

Table 2.1: Output of pytest Unit Tests and results for CT Analysing Library

Result	Test	
Passed	CTData.py::test_aggregate_spike_averages	
Passed	d CTData.py::test_clean_data_maximum_removed	
Passed	CTData.py::test_clean_data_minimum_removed	
Passed	CTData.py::test_load_additional_data	
Passed	CTData.py::test_load_additional_data_no_data	
Passed	CTData.py::test_load_data	
Passed	${\bf CTData.py::} {\bf test_NoDataFoundException}$	
Passed	Data_transforms.py::test_box_cox_data	
Passed	Data_transforms.py::test_pca_to_table	
Passed		
Passed	Data_transforms.py::test_standardise_data	
Passed	Graphing.py::test_plot_boxplot_as_dataframe	
Passed	Graphing.py::test_plot_boxplot_as_object	
Passed	Graphing.py::test_plot_difference_of_means	
Passed	Graphing.py::test_plot_histogram_as_dataframe	
Passed	Graphing.py::test_plot_histogram_as_object	
Passed	d Graphing.py::test_plot_pca	
Passed	ed Graphing.py::test_plot_qqplot	
Passed	Statistical_tests.py::test_baysian_hypothesis_test	
Passed	Statistical_tests.py::test_t_test	
Passed	Statistical_tests.py::test_test_normality	

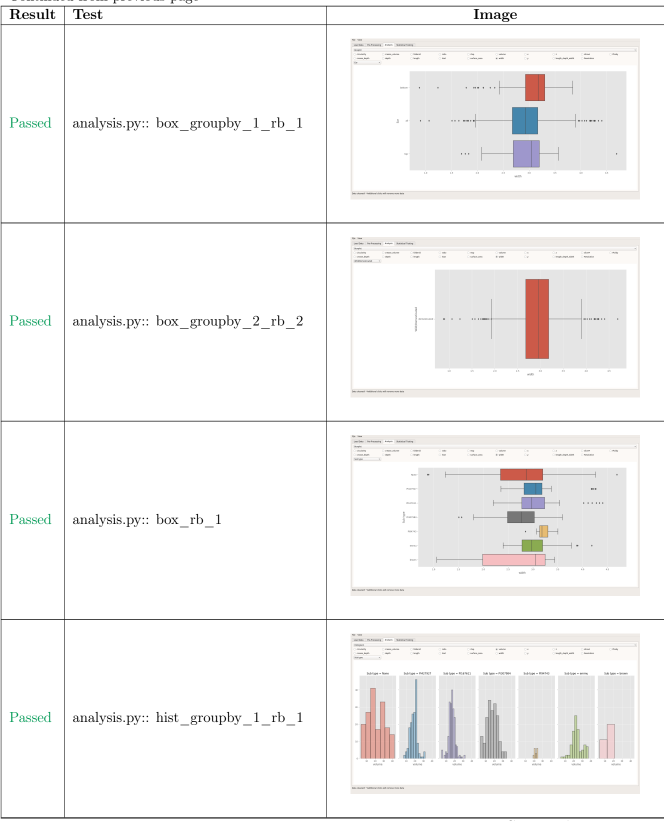
Unit Testing CT GUI Application

Table 2.2: Output of pytest Unit Tests and results for CT GUI Application

Result	Test	Image

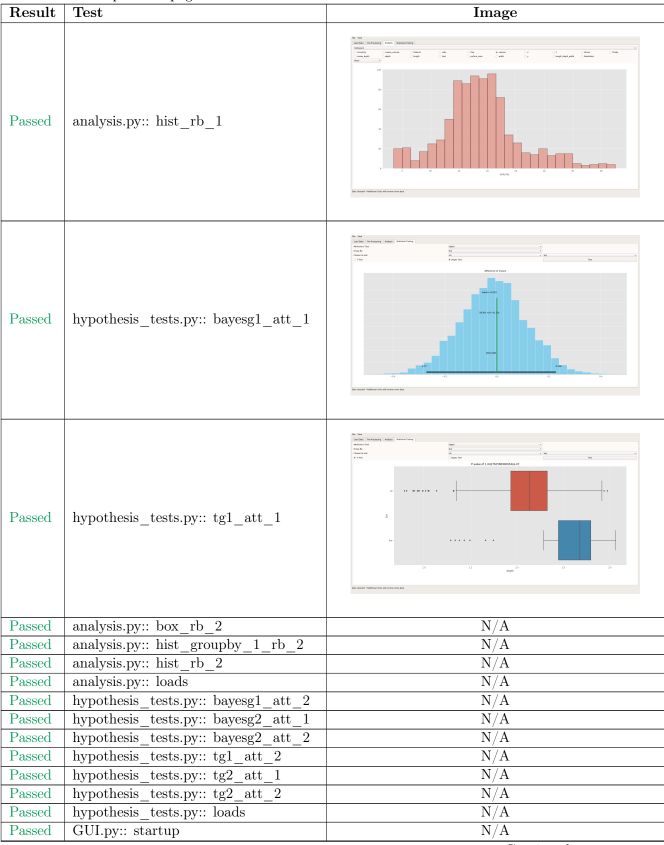
Continued on next page

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Result	Test	Image
Passed	load_data.py::	N/A
	load_data_with_rachis	
Passed	load_data.py::	N/A
	$load_data_without_rachis$	
Passed	preprocessing.py::	N/A
	$clean_data_remove_large$	
Passed	preprocessing.py::	N/A
	clean_data_remove_none	
Passed	preprocessing.py::	N/A
	$clean_data_remove_small$	
Passed	preprocessing.py::	N/A
	clean_data_remove_small_and_large	
Passed	preprocessing.py::	N/A
	load_additional_data	
Passed	preprocessing.py::	N/A
	load_additional_data_expected_fail	

Methods and Solutions

Data Pipeline

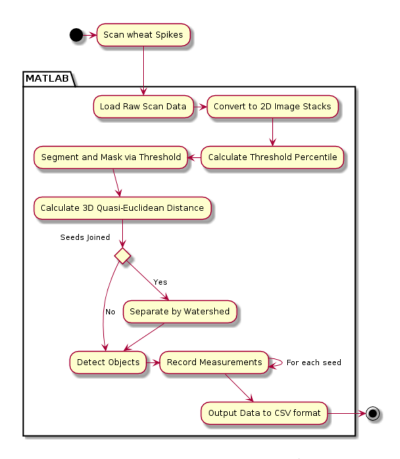


Figure 3.1: Image Processing Pipeline

Image Analysis Methods

New Watershed Algorithm

In order to solve the problem of misidentified and joint seeds, from the primitive collection, a *quasi-euclidean* distance transform was implemented into the analysis pipeline (figure:). This provided much better results than the previous *chessboard* transform which had been successful on more uniform data in previous studies [1].

Quasi-Euclidean algorithm

This algorithm measures the total euclidean distance along a set of horizontal, vertical and diagonal line segments [25].

$$|x_1 - x_2| + (\sqrt{2} - 1), |x_1 - x_2| > |y_1 - y_2| (\sqrt{2} - 1) |x_1 - x_2|, \text{ otherwise}$$
 (3.1)

In order to apply this to a 3D space Kleinberg's method is used [26]. This allows for nearest neighbour pixels to be sorted by \$k\$-dimensional trees and enabling fast distance transforms via Rosenfeld and Pfaltz's quasi-euclidean method stated in equation:3.1.

Effect of Enhanced Watershed algorithm

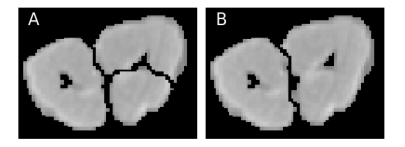


Figure 3.2: A showing the chessboard method, B improved quasi-euclidean method

Extracted Grains

CT Analysing Library Methods

CT GUI Application Methods

Data Analysis Methods

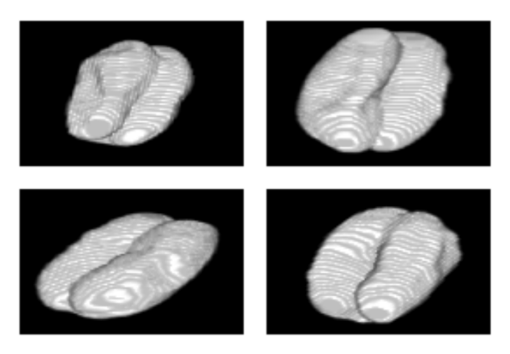


Figure 3.3: Individual Wheat grains, rendered in 3D

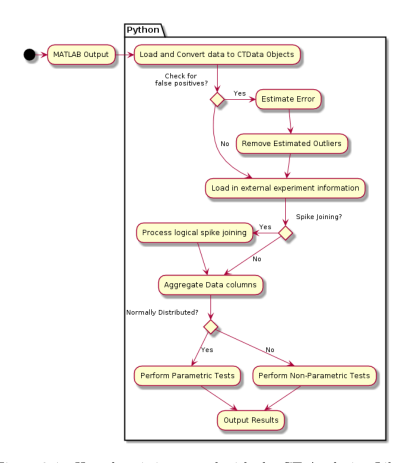


Figure 3.4: How data is integrated with the CT Analysing Library

Results

Discussion

Similar Research

Alternate Solutions

Critical Evaluation

Organisational Methods

Relevance to Degree

Time Management

Collaborative Work

Other Issues

Appendix

Software Packages Used

Libraries

Table 7.1: Software libraries used

MATLAB Image Processing	Numpy	Matplotlib
Toolbox		
Seaborn	Scipy	Sklearn
Statsmodels	Pymc3	Xlrd
PyQt5	gcc	

Tools

Table 7.2: Software tools used

MATLAB	Python Debugger (PDB)	IPython
Emacs	git	org-mode
Tomviz	ImageJ	

Glossary

Wheat Varieties

Code Segments and Examples

MATLAB Watershedding

Custom Documentation Generator

Table 7.3: Dictionary for Terms and acronyms

Term	Definition
р-СТ	Micro Computed Tomography
Genotype	A genetically distinct individual or group
Phenotype	A physical/measurable trait
Alleles	A variant of a gene
Genus	Classification ranking, below the family grouping
Genome	The complete genetic make up of an organism, which defines its individuality
Morphometric	The shape and form of an organism
GUI	Graphical User Interface
PCA	Principal Component Analysis
Spike	A singular stalk of wheat
Spikelet	A group of seeds all forming from the same node in a spike
MVC	Model View Controller - A design pattern for GUIs

Table 7.4: Dictionary for Wheat names used

Used name	Species name	
Monococcum	Triticum monococcum	
Monococcum Wild	Triticum Boeticum	
Tauschii	Aegilops tauschii	
Durum	Triticum Durum	
Dicoccoides	Triticum Dicoccoides	
Dicoccum	Triticum Dicoccum	
Ispahanicum	Triticum Ispahanicum	
Timopheevii	Triticum Timopheevii	
Spelta	Triticum Spelta	
Aestivum	Triticum Aestivum	
Compactum	Triticum Compactum	

```
function [W] = watershedSplit3D(A)
  % Takes image stack A and splits it into stack W
  % Convert to BW
 bw = logical(A);
  % Create variable for opening and closing
 se = strel('disk', 5);
  % Minimise object missshapen-ness
 bw = imerode(bw, se);
 bw = imdilate(bw, se);
  % Fill in any left over holes
 bw = imfill(bw,4,'holes');
  % Use chessboard for distance calculation for more refined splitting
 chessboard = -bwdist(~bw, 'quasi-euclidean');
  % Modify the intensity of our bwdist to produce chessboard2
 mask = imextendedmin(chessboard, 2);
 chessboard2 = imimposemin(chessboard, mask);
  % Calculate watershed based on the modified chessboard
 Ld2 = watershed(chessboard2);
  % Take original image and add on the lines calculated for splitting
 W = A;
 W(Ld2 == 0) = 0;
end
```

Listing 1: MATLAB Watershedding function

```
(defun populate-org-buffer (buffer filename root)
  (goto-char (point-min))
  (let ((to-insert (concat "* " (replace-regexp-in-string root "" filename) "\n") ))
    (while (re-search-forward
            (rx (group (or "def" "class"))
                space
                (group (+ (not (any "()"))))
                (? "(" (* nonl) "):" (+ "\n") (+ space)
                   (= 3 "\"")
                   (group (+? anything))
                   (= 3 "\"")))
            nil 'noerror)
      (setq to-insert
            (concat
             to-insert
             (if (string= "class" (match-string 1))
                 "** "
               "*** ")
             (match-string 2)
             "\n"
             (and (match-string 3)
                  (concat (match-string 3) "\n"))))helm-semantic-or-imenu
    (with-current-buffer buffer
      (insert to-insert))))
(defun org-documentation-from-dir (&optional dir)
  (interactive)
  (let* ((dir (or dir (read-directory-name "Choose base directory: ")))
         (files (directory-files-recursively dir "\py$"))
         (doc-buf (get-buffer-create "org-docs")))
    (dolist (file files)
      (with-temp-buffer
        (insert-file-contents file)
        (populate-org-buffer doc-buf file dir)))
    (with-current-buffer doc-buf
      (org-mode))))
```

Listing 2: Custom lisp code for generating easy to read documentation

Self-Documenting Code Example

```
def get_spike_info(self, excel_file, join_column='Folder#'):
    This function should do something akin to adding additional
    information to the data frame
    Onote there is some confusion in the NPPC about whether to use
    folder name or file name as the unique id when this is made into
    end-user software, a toggle should be added to allow this
    Oparam excel_file a file to attach and read data from
    Oparam join_column if the column for joining data is
    different then it should be stated
    try:
        # Grab the linking excel file
       info = pd.read_excel(excel_file,
                             index_col='Folder#')
       features = list(info.columns)
        # Lambda to look up the feature in excel spreadsheet
       def look_up(x, y): return info.loc[x['folderid']][y]
        # Lambda form a series (data row) and apply it to dataframe
       def gather_data(x): return pd.Series(
            [look_up(x, y) for y in features])
       self.df[features] = self.df.apply(gather_data, axis=1)
    except KeyError as e:
       print('Error matching data')
       print(e)
       raise NoDataFoundException
   except AttributeError as e:
       print(e)
       raise NoDataFoundException
```

Listing 3: Example of code documentation and readability from data transforms.py

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