Assembly of a Complex Genome: Defining Elements of Structure and Function

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Abstract

The post-human genome sequencing project era has seen an influx of genome sequencing projects established to investigate the structure, composition and characteristics of plant genomes. While the genome sequences of smaller plant genomes (ie. Rice) are currently available, there has been a lack of progress on the study of large, complex genomes such as barley (*Hordeum vulgare*) and wheat (*Triticum aestivum*), due to the difficulties in their sequencing and assembly. The aim of this study is to assemble and annotate targeted regions of chromosome 3B from *Triticum aestivum* cv. Chinese Spring (CS) and Hope. This study also aimed to complete a comprehensive, inter- and intra-species comparative analysis using Bioinformatics tools and strategies, in order to define structural and functional elements within the genome.

Genome sequences totalling 2.7Mb from two different loci of chromosome 3B in two different cultivars (*ctg11* from the short arm of CS, *ctg1034* from the long arm of CS and three assembled sequences over the equivalent *ctg11* region of Hope) were assembled using a novel ‘two-phase’ process that integrated information from a genome sequence assembler and a Triticeae-specific transposable element database. Through comparative genomics analysis a gene island was identified within a highly repetitive, heterochromatic region on 3BL that was highly conserved over four other cereal genomes (*Brachypodium distachyon*, *Oryza sativa*, *Sorghum bicolor* and *Zea mays*). Chromodomain-containing long terminal repeats from the *gypsy* family of retrotransposons were identified adjacent to the gene island and may suggest an involvement in the targeted insertion of transposable elements at the loci, protecting the gene-island from
dynamic evolutionary change. Characterisation of the \textit{ctg11} (\textit{Sr2} region) genome sequence on 3BS, identified a large \textasciitilde60kb mitochondrial genome insert and three members of the multi-gene beta-expansin family, with sequence analysis indicating local duplication within the sequence and rearrangements when compared to the equivalent region in a different wheat cultivar. \textit{In silico} and real-time transcription analysis of the individual gene was also confirmed. Within the equivalent \textit{ctg11} in Hope, a germin-like protein (GLP) cluster was identified and characterised that distinguishes between the two wheat cultivars. The genes in this GLP cluster were identified to belong to a sub-gene family that conferred broad level basal resistance in transient over-expressed systems in rice and barley.

The main outcome of this study was the development of a novel strategy of genome sequence assembly by utilising the complex component of the wheat genome that made assembly difficult: transposable elements. The complex genome sequence assembly methodology outlined in this thesis is suitable to be used as a model for future sequence assembly studies. The assembly of large pseudomolecule sequences (among the largest and most complete ever assembled in the wheat genome) enabled the Bioinformatics analysis of a representative sample of wheat chromosome 3B, providing valuable \textit{in silico} outputs for future functional analyses and allowing an in-depth intra- and inter-species comparative analysis with related genomes.
Declaration

Except where otherwise indicated, all work in this thesis is based on work carried out at the Centre for Comparative Genomics (CCG), Murdoch University, Australia. I declare that this thesis is my own account of my research and contains as its main content work, which has not previously been submitted for a degree at any tertiary education institution.

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James Breen
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My supervisors Professor Matthew Bellgard and Professor Rudi Appels deserve a lot of credit for this thesis. I thank Rudi for his enthusiasm, persistence, gentle encouragement and helpful advice, not only in this project, but also in my pursuit of future employment in genome research. I thank Matthew for taking a chance on me as an undergraduate student four years ago and having faith in my research.

Personally I would like to thank my parents, in-laws, grandparents, friends and extended family for their constant support. While they may not have understood much of my work, they were always excited and proud whenever I spoke about interesting developments. Special thanks also goes to my grandfather Professor Valentine Pervan for inspiring me to achieve excellence in academia. The high standards that he has set throughout his life is what I strive to reach.
Lastly I would like to thank my wife Nadine. Words cannot express the gratitude and appreciation I have for her support over the last three years. This thesis would not be possible without her. I dedicate this thesis to her.
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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>BAC</td>
<td>Bacterial Artificial Chromosome</td>
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<tr>
<td>BLAST</td>
<td>Basic Local Alignment Search Tool</td>
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<td>CCG</td>
<td>Centre for Comparative Genomics</td>
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<tr>
<td>CDD</td>
<td>NCBI Conserved Domains Database</td>
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<td>CS</td>
<td>Chinese Spring</td>
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<td>CSIRO</td>
<td>Commonwealth Scientific and Industrial Research Organisation</td>
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<td>CSRDB</td>
<td>Cereal smRNA Database</td>
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<td>DDBJ</td>
<td>DNA Databank of Japan</td>
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<tr>
<td>EBI</td>
<td>European Bioinformatics Institute</td>
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<td>EMBL</td>
<td>European Molecular Biology Laboratories</td>
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<tr>
<td>EMBOSS</td>
<td>European Molecular Biology Open Software Suite</td>
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<tr>
<td>EST</td>
<td>Expressed Sequence Tag</td>
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<tr>
<td>FPC</td>
<td>Fingerprinted Contig</td>
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<tr>
<td>GBrowse</td>
<td>Generic Genome Browser</td>
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<tr>
<td>GFF</td>
<td>Generic Feature Format</td>
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<td>GLP</td>
<td>Germin-like Protein</td>
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<td>GRR</td>
<td>Gene-Rich Region</td>
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<td>GyDB</td>
<td>Gypsy Mobile Element Database</td>
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<tr>
<td>HMM</td>
<td>Hidden Markov Model</td>
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<tr>
<td>InDel</td>
<td>Insertion/Deletion</td>
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<tr>
<td>ISBP</td>
<td>Insertion-Site Based Polymorphism</td>
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<tr>
<td>Acronym</td>
<td>Full Form</td>
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<tr>
<td>INRA</td>
<td>Institut National de la Recherche Agronomique (France)</td>
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<tr>
<td>IWGSC</td>
<td>International Wheat Genome Sequencing Consortium</td>
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<tr>
<td>LINE</td>
<td>Long Interspersed Nuclear Element</td>
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<tr>
<td>LTR</td>
<td>Long Terminal Repeat</td>
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<td>MITE</td>
<td>Transposable Element in Miniature</td>
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<td>MSU</td>
<td>Michigan State University</td>
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<tr>
<td>MTP</td>
<td>Minimum Tiling Path</td>
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<tr>
<td>MULE</td>
<td>Mutator-like Transposable Element</td>
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<tr>
<td>MYA</td>
<td>Million Years Ago</td>
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<tr>
<td>NCBI</td>
<td>National Centre for Biotechnology Information</td>
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<tr>
<td>NUMT</td>
<td>Nuclear Mitochondrial Genomic Insert</td>
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<tr>
<td>ORF</td>
<td>Open Reading Frame</td>
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<tr>
<td>OXOX</td>
<td>Oxalate Oxidase</td>
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<td>PCR</td>
<td>Polymerase Chain Reaction</td>
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<tr>
<td>PDB</td>
<td>Protein Databank</td>
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<td>PFAM</td>
<td>Protein Family Database</td>
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<td>QTL</td>
<td>Quantitative Trait Loci</td>
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<tr>
<td>RT-PCR</td>
<td>Real-Time Polymerase Chain Reaction</td>
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<tr>
<td>SINE</td>
<td>Short Interspersed Nuclear Element</td>
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<tr>
<td>SOD</td>
<td>Superoxidase Demutase</td>
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<td>SOE</td>
<td>Son of Eric</td>
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<td>TE</td>
<td>Transposable Element</td>
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<td>Acronym</td>
<td>Description</td>
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<tr>
<td>TIGR</td>
<td>The Institute of Genomic Research</td>
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<tr>
<td>TIR</td>
<td>Terminal Inverted Repeat</td>
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<tr>
<td>TREP</td>
<td>Triticeae Repeat Element Database</td>
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<tr>
<td>TSD</td>
<td>Target Site Duplication</td>
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<tr>
<td>UTR</td>
<td>Untranslated Region</td>
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<td>WA</td>
<td>Western Australia</td>
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General Introduction and Chapter 1

**Figure 1.1:** The conventional genome sequence assembly and annotation methodology employed in this study (on the left hand side). On the right hand side of the figure are general thesis questions investigating whether or not this conventional methodology is applicable to complex genomes such as hexaploid wheat.

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**Chapter 3**

**Figure 3.1:** Location of ctg1034. The bin 3BL-7 is characterised by the SSRs Xgwm299, Xgwm2152, Xgwm547, Xgwm247, Xgwm340, Xgwm181, Xfwm4, Xcfa2170, Xbarc84, Xpsr170, XksuG62 (Sourdille et al. 2004) and the two ISBP markers (Sc3-119 and Sc3-120) could be assigned to this deletion bin. Lane 1-8 on the electrophoresis gel on the left-hand panel of the figure indicates the analysis of Sc3-119 and Sc3-120 with the deletion bins of chromosome 3B (lane 1: 3BS-8, lane 2: 3BS-9, lane 3: 3BS-1, lane 4: 3BL-10, lane 5: 3BL-7, lane 6: Halberd and lane 7: Cranbrook). Genetic mapping using the Cranbrook x Halberd population (McFadden et al. 2007) confirmed the long arm location on 3B. The genetic map (middle panel) shows a small part of the map (complete map, Cran*Hal 3B Feb09, is available at http://ccg.murdoch.edu.au/cmap/ccg-live/).

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Chapter 5

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**Chapter 6**

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(* Products showing poor quality sequence repeated in Table 2.9).

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(* Products showing poor quality sequence repeated in Table 2.9).

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General Introduction

Wheat is an important world food crop contributing >60% of the total calories consumed in the world daily. The temperate environment that wheat is able to be grown in means that it takes up more area of land that any other agricultural crop (Gill et al. 2004). In order to feed an exponentially increasing world population, crop yields need to be improved and research must be carried out to protect crops from abiotic-stress (extreme environmental conditions such as drought) and biotic-stress (attack from fungi, parasites and viruses that cause plant diseases).

Genome sequencing, enabling the identification of important agronomic genes, is an important step in improving crop characteristics and yields. Research into plant species such as *Oryza sativa* (rice), *Sorghum bicolor* and *Arabidopsis thaliana* have all benefited from a fully sequenced genome and their small and compact genomes allowed them to be sequenced relatively quickly and easily. Sequencing the wheat genome on the other hand has been impeded by its extremely large genome size (~16Gb; 37 times larger than the entire rice genome sequence), hexaploid genome structure and its high complexity (due a high proportion of repetitive sequences caused by transposable elements (TEs)).