Scalable and reproducible genome analysis in the age of next-generation genome sequencing

by

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TABLE OF CONTENTS

ABSTRACT	vi
CHAPTER 1. INTRODUCTION	1
Overview	1
Polistes dominula genome project	2
ParsEval: comparison of distinct annotation sources	4
iLoci: an organizational framework	5
Dissertation Organization	6
CHAPTER 2. GENOME, TRANSCRIPTOME, AND METHYLOME	
SEQUENCING OF A PRIMITIVELY EUSOCIAL WASP REVEAL	R 1. INTRODUCTION
A GREATLY REDUCED DNA METHYLATION SYSTEM IN A	INTRODUCTION 1
SOCIAL INSECT	7
Abstract	7
Introduction	8
Materials and Methods	11
Sample collection and sequencing	11
Genome assembly and annotation	12
Transcriptome assembly and annotation	12
	12
Methylome analysis	13
Examination of interfamilial relationships via gene tree analysis	13

Results and Discussion	14
Genome description and assessment of de novo genome quality	14
Composition of P. dominula genome shows a combination of typical hy-	
menopteran as well as unique features	16
Caste-related transcriptome reveals differentially expressed conserved and	
novel genes	18
DNA methylation system is greatly reduced in <i>P. dominula</i>	21
Aculeate Phylogeny.	26
Conclusions	27
Acknowledgments	28
Data Accessibility	29
Author Contributions	29
Figures	30
CHAPTER 3. PARSEVAL: PARALLEL COMPARISON AND ANAL-	
YSIS OF GENE STRUCTURE ANNOTATIONS	36
Abstract	36
Background	37
Implementation	40
Overview	40
Gene locus identification	41
Gene structure representation	42
Comparative analysis of annotations	43
Reporting comparison scores	45
Results and Discussion	46
Use case: predictions vs. gold standard	47
Use case: two sets of annotations	48
Benchmarks	50

Performance evaluation in comparison to Eval software	53
Conclusions	54
Availability and requirements	55
Authors contributions	55
Acknowledgements	55
Figures	56
CHAPTER 4. ILOCI: ROBUST GENOME ANNOTATION AND ANAI	
YSIS FOR PROVISIONAL GENOME ASSEMBLIES	59
Abstract	59
Background	60
Methods	62
Toolkit design	62
Conceptual definition of interval loci	64
Operational definition of iLoci	65
Implementation	68
Genome content statistics	69
Genome organization statistics	70
Comparing assembly/annotation pairs: iLocus stability	72
Comparing genome content and organization between related genomes:	
homologous iLoci	72
Data sets analyzed	73
Results	75
iLoci provide an informative decomposition of genome content	75
iLoci reflect patterns of genome organization	77
Characterization of three species of interest	81
iLoci provide a robust representation of the genome	83
iLoci reveal extent of gene conservation in green algae	86

Discussion	87
Conclusions	89
Figures	90
CHAPTER 5. CONCLUSIONS	102
BIBLIOGRAPHY 1	106
ACKNOWLEDGEMENTS	119

ABSTRACT

Recent advances in DNA sequencing technology and a proliferation of new algorithms for assembling, annotating, and analyzing genomes have made genome-scale sequencing more accessible than ever. As a result, the last several years have seen a dramatic increase in the number of published draft genomes. Many important research problems revolve around interpretation of these draft genomes: What are the contents of a genome? How many genes are there? Are there any conspicuous losses of genes of interest? Is the genome compact, with genes clustered very tightly, or are genes separated by large intergenic spaces? Are intergenic spaces distributed evenly throughout the genome? Which characteristics of genome composition and organization are well conserved, and which appear to be unique, warranting further investigation?

In this dissertation, I investigate this topic in multiple contexts. First, I present a draft genome of the paper wasp *Polistes dominula*, a model species for study of the evolution of social behavior. The genome of *Polistes* is similar to other social insects in many respects, but has an extremely biased nucleotide composition and shows some evidence of a reduction in genome size. Analysis of transcriptome and methylome data from queen and worker wasps reveals evidence of caste-related differences in gene expression, as well as a tremendous reduction in DNA methylation, previously thought to be an important factor in caste differentiation.

Second, I investigate questions of genome composition and organization more generally. Given a new genome assembly and annotation, what can we determine quickly about the genome's contents? What can be said about the distribution of genes and the overall "compactness" of the genome? How should this be compared to previously pub-

lished results for related species? I present a framework (and related tools) that provides precise solutions to these questions, and discuss insights gained by applying these tools to study various model organism genomes.

CHAPTER 1. INTRODUCTION

Overview

In the 2000s, the advent of new nucleotide sequencing strategies based on ion semi-conductors (Ion Torrent), pyrosequencing (454), and sequencing-by-synthesis (Illumina) provided new tools for studying genomes of both model and non-model organisms at unprecedented scale, resolution, and cost effectiveness. These technologies continue to evolve, more recent innovations involving single-molecule long read sequencing (Pacific Biosciences SMRT and Oxford Nanopore). By the 2010s, these so-called next-generation sequencing (NGS) technologies had made genome sequencing accessible to essentially any scientist with even a modest research budget. This "democratization of sequencing" has precipitated a tremendous increase in the number of published genome projects and draft genome sequences (http://www.ncbi.nlm.nih.gov/genome/browse/), as well as genome-scale data sets profiling gene expression, chromatin accessibility, transcription initiation, and a multitude of other genomic characteristics.

During this same time frame, however, the democratization of genome analysis has occurred to a much lesser extent. The availability of high-quality model reference genomes has changed very little. And as newly acquired data continues to flood into public databases, many scientists struggle to effectively manage the data and critically evaluate downstream research products. The proliferation of new algorithms and software tools for analyzing NGS data is a mixed blessing for scientists who now have both the flexibility and the burden of selecting suitable tool(s) for a particular analysis.

The complexity and difficulty of genome assembly [1, 2, 3, 4], annotation [5, 6, 7], and analysis [8, 9, 10] has been reported in various recent studies and community projects. The consistent, resounding message from this growing body of work is that genomics data quality varies considerably across data sets (and even within a single data set), and that the performance of state-of-the-art algorithms is difficult to predict on new data. Without a well-funded and well-staffed research consortium to manage the painstaking work of gap-filling each new genome assembly and carefully curating its contents, the new reality is that most reference genome assemblies will remain fragmented and unfinished, and that provisionally annotated genome features will fall on a wide spectrum of reliability. Doing principled and reproducible science in this setting requires disciplined quality control and data evaluation. The focus of this dissertation has been the development of a framework—and associated software tools—to enable robust annotation and analysis of NGS-based genomics data, as motivated by research problems I encountered in genomics research projects.

Polistes dominula genome project

My first encounter with many of these issues came from my genomics studies of the paper wasp *Polistes dominula*. This wasp is an important model system for studying the evolution of social behavior, as it exhibits an intermediate level of social complexity, with no morphological differentiation between castes and frequent competition for reproductive opportunities within colonies [11]. The molecular mechanisms underlying caste differentiation across all social insects are still poorly understood, although various mechanisms have been proposed to play primary roles, such as differential gene expression [12, 13, 14, 15, 16, 17, 18], differential splicing [19], DNA methylation patterns [19, 20], and recently emerged species- or clade-specific genes [21, 22]. These hypotheses have been tested in bees and ants, but prior to this study no genome resources were avail-

able for any species from the major vespid wasp lineage. The *Polistes dominula* genome project was funded to sequence the genome, transcriptome, and methylome of the wasp, to facilitate investigation of the molecular basis of caste differentiation in *Polistes*, and to provide an additional important data source for comparative analysis of all social insects.

Our initial work was driven largely by questions of genome composition. How large is the *Polistes* genome? What is its nucleotide composition? How many genes does it encode? Are the small handful of well-known "social behavior" genes present in the genome as expected? In short, is there anything that immediately distinguishes the paper wasp genome from genomes of related species?

The next set of questions were driven by a comparative genomics perspective. How does the size and composition of the *Polistes* genome compare to the bees and the ants? What proportion of annotated gene models are well-conserved within the Hymenoptera? Can conserved single-copy orthologs provide any insight into the unresolved evolutionary lineage of the bees, ants, and wasps?

At the same time, we investigated questions of functional genomics and epigenomics. How many *Polistes* genes show differential expression between the queen and worker castes? What is the extent of alternative splicing in *Polistes*, and do any alternative splicing events exhibit caste-related bias? What is the extent of DNA methylation in *Polistes*, and does the genome encode a full complement of methylation-related genes? This last question led us to one of the highlight discoveries of the study: that *Polistes* lacks a critical DNA methylation genome-wide.

More in line with the focus of this dissertation, the *Polistes dominula* genome project exposed me to the challenges of creating genomic data resources *de novo* for a non-model research system, and the corresponding challenges of data quality assessment and management. These challenges motivated the development of methods and tools to facilitate comparison, evaluation, and analysis of genome annotations, with additional

applications to studying genome organization.

ParsEval: comparison of distinct annotation sources

Automated genome annotation typically relies on integrating tools for ab initio gene prediction, transcript and protein spliced alignment, and evaluation of support for gene structural components. Each class of tools comes with a variety of parameter settings, and it can be difficult to predict in advance the influence these parameters will have on the final annotation product. When annotating a non-model genome de novo, it is often necessary to refine parameter selection on a small subset of the available data before proceeding to annotate the entire genome. Subsequent re-annotation is often necessary when additional data (such as ESTs or RNA-Seq reads) become available, or when improved gene prediction methods are published. And in some cases, the scientists annotating a particular genome have a vested interest in improving the accuracy of the annotation software itself. In each of these scenarios, a primary objective is to identify similarities and differences between annotations derived from different parameter settings or alternative workflows, to facilitate evaluation of the annotations.

Development of a new annotation pipeline (CpGAT) within our research group provided the initial motivation for ParsEval. Evaluating this tool's performance in comparison to a gold-standard annotation was impractical manually, and existing software tools [23, 24] lacked important features such as locus-scale resolution. I created the ParsEval program to address the need for genome-scale evaluations with locus-scale detail. Later during the preliminary stages of the *Polistes dominula* genome project, ParsEval proved a valuable data assessment tool as we tuned our genome annotation workflow.

The ParsEval paper introduces a precise operational definition for a *gene locus* based on the locations of annotated gene models. The objective was to define a parsing of the genome into distinct units that are complete and can be analyzed independently. Gen-

eralizing this concept and applying it as an organizational principle for genome analyses has been a major focus of this dissertation.

iLoci: an organizational framework

The ParsEval tool proved valuable throughout the *Polistes dominula* genome project, but we subsequently encountered issues that required additional attention and development. Although our *Polistes* assembly and annotation compared favorably to other published hymenopteran genomes, our data was subject to the same quality inconsistencies that characterize any NGS-based genome project. Answering questions related to gene expression and genome composition therefore required careful consideration of, for example, how precisely to handle overlapping gene models and how to distinguish differences rooted in biology from technical artifacts. It was in addressing these issues that we extended the *gene locus* definition introduced by ParsEval and developed the *interval locus* (*iLocus*) as a more generalized organizational framework for genome analyses.

iLoci define an unambiguous parsing of an annotated genome sequence into distinct regions, each encapsulating the genomic context of a gene or intergenic space. This parsing provides a complete and granular decomposition of the genome, and characteristics of iLoci are reflective of a variety of genomic features of interest. In parallel with the *P. dominula* genome project in which iLoci played a prominent role, we investigated general applications of iLoci. We discuss their utility for describing the gene content of a genome and for applying quality control when calculating diagnostic characteristics of a genome. We demonstrate the utility of iLoci for characterizing genome organization, within a single genome and between multiple genomes in a clade of species. Finally, we report the stability of iLoci across distinct assembly and annotation versions, highlighting their utility as reproducible units of analysis.

Dissertation Organization

This dissertation is organized into five chapters. Chapter 1 provides an overview of the dissertation, a motivation for the work, and a brief discussion of relevant literature. Chapters 2 through 4 are presented as self-contained manuscripts: chapter 2 is a research paper published in *Molecular Ecology* describing the genome, transcriptome, and methylome of the paper wasp *Polistes dominula*, highlighting its reduced DNA methylation system, several hundred loci with caste-related differential expression, and the lack of any detectable caste-related differential splicing in the adult organism; chapter 3 is a paper published in *BMC Bioinformatics* describing *ParsEval*, a tool for comparing two alternate sources of annotation for a genome sequence; chapter 4 is a methodology paper to be submitted to *Genome Biology*, describing the use of *interval loci (iLoci)* as an organizational framework for reproducible genome analysis; Chapter 5 provides brief concluding remarks and suggestions for further research.

CHAPTER 2. GENOME, TRANSCRIPTOME, AND METHYLOME SEQUENCING OF A PRIMITIVELY EUSOCIAL WASP REVEAL A GREATLY REDUCED DNA METHYLATION SYSTEM IN A SOCIAL INSECT

A manuscript published in *Molecular Ecology*.

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Abstract

Comparative genomics of social insects has been intensely pursued in recent years with the goal of providing insights into the evolution of social behavior and its underlying genomic and epigenomic basis. However, the comparative approach has been hampered by a paucity of data on some of the most informative social forms (e.g. incipiently and primitively social) and taxa (especially members of the paper wasp family Vespidae) for studying social evolution. Here we provide a draft genome of the primitively eusocial model insect *Polistes dominula*, accompanied by analysis of caste-related transcriptome and methylome sequence data for adult queens and workers. *P. dominula* possesses a fairly typical hymenopteran genome, but shows very low genome-wide GC content and some evidence of reduced genome size. We found numerous caste-related differences in gene expression, with evidence that both conserved and novel genes are related to caste differences. Most strikingly, these –omics data reveal a major reduction in one of the

major epigenetic mechanisms that has been previously suggested to be important for caste differences in social insects: DNA methylation. Along with a conspicuous loss of a key gene associated with environmentally responsive DNA methylation (the de novo DNA methyltransferase Dnmt3), these wasps have greatly reduced genome-wide methylation to almost zero. In addition to providing a valuable resource for comparative analysis of social insect evolution, our integrative –omics data for this important behavioral and evolutionary model system call into question the general importance of DNA methylation in caste differences and evolution in social insects.

Introduction

The rapidly increasing availability of genomic resources for non-traditional model organisms with well-developed social behavior has incited great interest in the genomic basis of complex social life, or sociogenomics [25]. Sociogenomic studies on a variety of species, from rodents to fish to insects, have provided a wealth of information about the transcriptomic and genomic characters associated with different forms of derived social behavior, from affiliative behavior, to aggression, to division of labor within animal societies [26]. To date, however, these studies have focused on relatively few species separately, and current data have not allowed a comprehensive comparative and phylogenetic approach to understanding the genomic changes that accompany the evolution of sociality. As advances in sequencing technology have greatly facilitated the generation of genome-scale data for emerging model species [27], it is an exciting time to seek integration of genomic, transcriptomic, and epigenomic data from species at key transitional points in the evolution of sociality from solitary behavior.

The eusocial insects are one of the most important sociogenomic model groups, comprising a diverse and ecologically successful group of animals with a highly derived form of social behavior characterized by the presence of reproductive and non-reproductive castes [28]. Eusocial insects are excellent models for understanding the evolution of complexity as the switch from solitary to eusocial life marks one of the major transitions in evolution due to the shift from individual selection to colony level selection [29]. Currently, published genome sequences are available for three parasitic *Nasonia* wasps [30], which provide a solitary outgroup for all of the social Hymenoptera species, and 20 eusocial insect genomes—ten bees [31, 32, 33, 34] and nine ants [35, 36, 37, 38, 39, 40, 41], and very recently, one paper wasp [42]. These studies provide important baseline data on genomic characters associated with eusociality.

Importantly however, prior comparative sociogenomic analyses within the social insects have suffered from two major deficits. First, there have been relatively scant genomic resources available for one of the three major hymenopteran eusocial lineages, the paper wasp family Vespidae (Figure 2.1A). Although social wasps, bees, and ants evolved from a common ancestor over 100 million years ago, these societies have independently evolved many convergent features, including the presence of female castes in the form of queens and workers. Second, there have been relatively few genome sequences available for species in key transitional stages between solitary and eusocial forms (notably, there has been a large recent advance in this area [31, 32, 42, 33]). Here, we expand the potential for comparative genomics of eusocial Hymenoptera by describing the first complete genome sequence of *Polistes dominula*, a behavioral model species within the family Vespidae that exhibits an intermediate form of social behavior, making it highly informative for studying the evolution of sociality [43].

Polistes wasps form small "primitively eusocial" societies containing queens and altruistic workers, but unlike honey bees, their colonies are characterized by prominent conflict over reproduction [44]. Queens and workers engage in dominance interactions and there is constant competition between females for reproductive opportunities. In addition, Polistes have small colonies with a relatively small number of individuals, and colonies are started anew annually by founding queens [45]. These characteristics

have made *Polistes* one of the main systems for testing hypotheses about the evolution of altruistic behavior [46]. For example, studies of cooperation and conflict in small groups of *Polistes* wasps have provided some key tests of how genetic relatedness can facilitate cooperation [47]. In addition, observations of *Polistes* behavior led to new hypotheses about the evolution of altruistic behavior from maternal behavior [46], which have derived some support from transcriptomic studies [48, 49]. A genome sequence for Polistes dominula, the best-studied member of the model genus Polistes, greatly enhances our power to study the genetics of social behavior via comparative genomic and transcriptomic analyses, allowing for the identification of protein coding changes, regulatory regions, and epigenetic modifications associated with sociality. To facilitate future comparative analyses, we provide a high quality draft P. dominula genome and describe informative features of this genome in reference to other previously published bee, ant, and Nasonia wasp genomes. Our genome sequence was derived from an invasive population (from Pennsylvania, USA) of *Polistes dominula*, a temperate species that is native to Europe. Our genome represents the second published paper wasp genome, the first being the very recently published genome of the Neotropical paper wasp *Polistes* canadensis [42]. Although both species are primitively eusocial, they are not closely related (split between Old and New World *Polistes* at 10-80 million years [50]) and have several differences in their ecology and social biology. Thus, we provide comparisons between these congeners, confirming many conserved *Polistes* genome characteristics but also highlighting some conspicuous differences between the two paper wasp genomes.

One of the most active recent areas of research in insect sociogenomics centers on the role of epigenetics in the regulation and evolution of eusociality. Recent studies suggest epigenetic modifications to DNA are ubiquitous within the social Hymenoptera [51, 52], and furthermore, various authors have suggested that differential DNA methylation during larval development contributes to caste differential gene expression and alternative splicing [19, 53], and differences in caste-related phenotypes [54] in both honey bees and ants. We previously hypothesized that DNA methylation might also be important for caste differential expression and behavioral and physiological caste differences in primitively eusocial species such as *Polistes dominula* [52, 55]. However, recent studies suggest DNA methylation may be less important for primitively eusocial species, including *Polistes canadensis* [31, 42]. Therefore, we looked for evidence of a functional DNA methylation system by investigating the presence of a full complement of DNA methylation enzymes in the *Polistes dominula* genome. In addition, we performed RNA-sequencing and whole genome bisulfite sequencing (methylome sequencing) on a set of adult queen and worker samples in order to examine caste-associated differential expression and DNA methylation in the *Polistes dominula* genome. These experiments provide a valuable point of comparison to other social insects for assessing whether DNA methylation is a shared, general mechanism related to sociality in insects.

Materials and Methods

Detailed protocols are provided in SI Materials and Methods.

Sample collection and sequencing

Five Illumina paired-end whole genome shotgun libraries were prepared from a single pupal male collected from an invasive population in State College, Pennsylvania. The libraries, ranging in insert size from <200bp to 8Kbp (see **Table S1**), were sequenced on the Illumina HiSeq 2000 platform.

Same-nest pairs of six adult workers and six egg-laying adult queens and from six different colonies were collected for transcriptome and methylome sequencing; these twelve individuals were from the same State College, Pennsylvania population as the pupal male used for genome sequencing. The head of each adult was cut in half, with RNA extracted from one half for transcriptome sequencing and DNA extracted from the other half for methylome sequencing.

Genome assembly and annotation

Five whole genome shotgun libraries were assembled using the AllPaths-LG genome assembler [56], with the smallest library designated as the fragment library and the other four libraries designated as jumping libraries. The assembled scaffolds were then screened for repetitive DNA, masked, and annotated by the MAKER pipeline [57]. The annotation workflow incorporated evidence from spliced alignments of transcripts from three Polistes species, spliced alignments of reference proteins from Apis mellifera and Drosophila melanogaster, gene models produced by three ab initio gene predictors, and manual gene annotations contributed via the PdomGDB genome browser's community annotation portal.

Transcriptome assembly and annotation

Twelve Illumina paired-end RNA-Seq libraries were assembled using the Trinity assembler [58] with the --CuffFly algorithm and the --jaccard_clip settings enabled. Assembled transcripts were then post-processed to discard contaminants, split transcript chimeras, and annotate transcript functions by similarity to known proteins and miRNAs. Two previously published *Polistes* transcriptomes [59, 18] were processed with the same procedure.

Differential expression analysis

Twelve Illumina paired-end RNA-Seq libraries, six from queens and six from workers, were sequenced and reads were mapped individually to the genome using Bowtie [60]. Preliminary examination of the alignments revealed an extremely wide dynamic range of expression values (<10 reads to millions of reads mapped per replicate) and in some cases considerable variation between replicates. To account for these observations we

discarded loci with too many or too few reads mapped (normalized by sequence length) or with a high coefficient of variation.

Expression was quantified using RSEM [61], and the EBSeq package [62] was used for identifying loci with caste differential expression at a false discovery rate of < 0.05. A complete description of the number of raw reads, data filtering procedure, and software parameters used is available in SI Methods and Results.

Methylome analysis

Two DNA samples (one pooled sample from workers and one pooled sample from queens, derived from the same six individuals used for transcriptome sequencing) were subjected to bisulfite treatment, and each sample was used to generate separate Illumina libraries for sequencing. The Bismark software [63] was used for read mapping and methylation calls. Highly supported methylation sites were determined as sites with significant number of methylation calls based on a binomial probability model with Bonferroni correction at the 1% significance level (assuming a 99.5% conversion rate in the treatment). We also reanalyzed existing datasets from from *Polistes canadensis* [42], honey bees [53], the ants *Harpegnathos saltator* and *Camponotus floridanus* [64], and the parasitoid wasp *Nasonia vitripennis* [65] in order to compare our *P. dominula* results to previously analyzed Hymenoptera (see SI Methods). Numbers of methylation sites were determined with the BWASP workflow (http://brendelgroup.github.io/BWASP/) using pooled reads from published data sets for each species and caste. In-depth descriptions of the comparative data, analysis pipelines, and comparative results are provided in a companion paper (Toth AL, Sankaranarayanan S, and Brendel VP, *in preparation*).

Examination of interfamilial relationships via gene tree analysis

Genes with conserved single-copy orthologs in *Apis mellifera*, *Harpegnathos saltator*, *Polistes dominula*, and *Nasonia vitripennis* were identified with protein clustering (see

SI Methods for clustering criteria). For each gene, a multiple sequence alignment of the four corresponding protein sequences was computed, and from that alignment a phylogenetic tree was inferred *via* maximum likelihood. After all gene trees had been constructed, each tree was analyzed to note its topology and collect a tally of the three possible topologies observed.

Results and Discussion

Genome description and assessment of de novo genome quality.

We used Illumina technology to sequence genomic DNA from a single, haploid pupal male P. dominula from an invasive population in State College, Pennsylvania, USA. The DNA was used to generate five Illumina genomic DNA libraries of varying insert size (Table S1), each sequenced on a single channel on an Illumina HiSeq instrument. This generated a total of 78.6 Gb of raw sequence, which was filtered using Trimmomatic [66] to remove sequencing adapters and low quality base calls. The groomed data were assembled using AllPaths-LG [56], producing 1,483 scaffolds with an N50 of 1.63 Mb and a combined length of approximately 208 Mb. The genomic reads provide approximately 319x coverage of the genome. This genome assembly compares favorably to Illumina-and 454-based assemblies of other social Hymenoptera, in particular to that of Polistes canadensis, which despite a higher level of fragmentation provides a consistent estimate of the Polistes genome size. Table 1 shows P. dominula in comparison to a few representative previously published paper wasp, bee, ant, and non-social wasp genomes (taxon selection described in SI Methods).

The genome assembly appears to be quite complete, on par with other Illuminabased draft insect genomes [35, 36]. CEGMA analysis [67] showed 246 (99.2%) of 248 ultra-conserved core eukaryote genes to be present in the genome assembly. The total assembled length of the genome approaches both *in silico* estimates of the *P. dominula*

Table 2.1 Genome assembly summary for $Polistes\ dominula$ and six other Hymenoptera.

Species	Clade	Sequencing strategy	Assembly level	Scaffolds	Total size	N50	Gene count	CEGMA genes recovered (%)
P. dominula	Paper wasps	Illumina	Scaffold	1483	208 026 220	1 625 592	12 153	246/248 (99.2)
Polistes canadensis	Paper wasps	Illumina	Scaffold	3836	211 209 432	521 566	15 755	245/248 (98.8)
Apis mellifera	Bees	Sanger/ SOLiD/ 454	Pseudochromosome	5645	250 270 657	13 219 345	13 275	239/248 (96.4)
Bombus terrestris	Bees	454	Pseudochromosome	5678	248 654 244	12 868 931	10 313	239/248 (96.4)
Harpegnathos saltator	Ants	Illumina	Scaffold	8893	294 465 601	601 965	12 254	239/248 (96.4)
Camponotus floridanus	Ants	Illumina	Scaffold	10 791	232 685 334	451 320	11 724	238/248 (96.0)
Nasonia vitripennis	Nonaculeate wasps	Sanger	Pseudochromosome	6169	295 780 872	897 131	13 794	234/248 (94.4)

genome size based on k-mer distributions in the sequence data (246 Mb, see **SI Results**) and earlier estimates based on flow cytometry (300 Mb, [68]). The gene space of the P-dominula genome therefore appears to be almost completely represented in the assembly, suggesting that the unrepresented portions of the genome are likely highly repetitive regions that are difficult to assemble with Illumina technology.

Automated annotation of *P. dominula* genes was based on a specifically trained MAKER workflow [57] and incorporated protein evidence from *Apis mellifera* (NCBI release 102 and OGS 3.2) and *Drosophila melanogaster* (FlyBase r5.55) and transcript data from *P. dominula* (described below), *P. metricus* [59], and *P. canadensis* [18]. Also integrated into the annotation were 180 gene models that, during preliminary stages of annotation, were manually curated and refined using the yrGATE portal [69]. This resulted in 11,819 predicted gene models, designated as release 1.2 (see **DATA ACCE-SIBILITY**). Similarity searches using BLAST revealed most of these predicted genes—10,755 out of 11,819 (91%)—have hits to the NCBI non-redundant database, whereas 1,064 show no significant similarity to known proteins. Of the genes with predicted homologs, most (10,504, or 98%) have best hits to other Hymenoptera annotated proteins (**Figure S3**). These gene models represent the first whole-genome annotation of a

vespid wasp, and include thousands of high-quality conserved genes enabling more detailed comparative analysis of hymenopteran genomes, as well as many species-specific gene models with which to investigate for evidence of novel clade-specific genes.

Composition of *P. dominula* genome shows a combination of typical hymenopteran as well as unique features.

Comparisons of the *P. dominula* genome assembly to those of other Hymenoptera revealed the assembled genome size and proportion of the genome occupied by transposable elements to be within the range of the other species. Published hymenopteran genomes show variety in the types and amounts of transposable element (TE) and other repetitive content, with *Apis mellifera* harboring almost exclusively a small number of *mariner* class transposons, and *Nasonia vitripennis* on the other hand harboring diverse repetitive elements constituting approximately a quarter of its genome [34, 30]. The *Polistes dominula* and *P. canadensis* genomes contain a fairly low level of repetitive DNA; 11-14% of the genome assemblies (24-30 Mb) are estimated to be repetitive, the majority of which represents simple repeats and low complexity sequence. The two *Polistes* genomes harbor a very similar cohort of TEs, dominated in both genomes primarily by L2/CR1/Rex and R1/LOA/Jockey LINEs, Gypsy/DIRS1 LTRs, and Tc1-IS630-Pogo DNA elements (Table S4).

Characteristics of genome structure were further investigated by parsing the *P. domin-ula* genome into 17,888 interval loci (iLoci), each iLocus capturing the local genomic context of a single gene (11,713 iLoci), a cluster of overlapping genes (205 iLoci), or an intergenic region (5,970 iLoci; see **SI Methods**). In order to compare a set of comparable genes across species and rule out differences due to annotation artifacts, homologous iLoci were determined by computing iLoci for several additional insect species and clustering their protein products. A comprehensive comparison included 17 hymenopteran species in total, but for illustrative purposes, seven representative species (*P. dominula*,

P. canadensis, two bees, two ants, and a non-aculelate, non-social hymenopteran out-group to the social insects, Nasonia vitripennis) are shown for comparison in this and subsequent analyses; see SI Methods.

At 11,918, the number of protein-coding iLoci (genes or gene clusters) in the P. dominula genome is well within the range observed in other Hymenoptera (see Figure S5). However, gene iLoci occupy only 73.0 Mb (35.1%) of the P. dominula genome (and a similar proportion was found in P. canadensis); this is much less compared to other species of Hymenoptera in which genes occupy between 140-160 Mb (and 50-65%) of the assembled genome (Figure 2.1B). This difference is due primarily to the annotation of fewer long genes, and in particular, long introns: while other Hymenoptera have 600-700 gene iLoci 50 kb in length or greater, P. dominula has only 84 (see Figure 2.1B). Further comparative genomic analyses can help resolve whether this observed reduction in long introns in the P. dominula genome is a truly unique characteristic of the genome sequence, or whether it stems from differences in annotation workflows.

The Polistes dominula genome is also characterized by an extremely biased nucleotide composition (Figure 2.1C). With a genome-wide GC content of 30.8%, the P. dominula genome is the most biased genome yet reported in Hymenoptera [35, 36, 38, 39, 40, 34, 30] and one of the most biased known in any animal (http://www.ncbi.nlm.nih.gov/genome/browse/). At the resolution of individual gene loci, however, P. dominula is not as GC-poor as Apis mellifera (29.0% and 24.7% median GC content, respectively), the primary factor being the extreme bias of introns in A. mellifera (21.3% median GC content for P. dominula versus 17.3% median GC content for A. mellifera; the distribution of intron length is nearly identical for P. dominula and A. mellifera). The composition of the P. canadensis genome is slightly less biased than P. dominula at all levels of resolution, but all trends in comparison to A. mellifera are consistent (see SI Results). The biased composition of Polistes genomes raises some compelling questions about the evolution of genome composition and potential contributing factors such as

bias in DNA mismatch repair and other genome maintenance mechanisms, as well as the possibility of historically high levels of CpG methylation and cytosine deamination.

Caste-related transcriptome reveals differentially expressed conserved and novel genes.

Distinct queen and worker castes arise from the same genome, a phenomenon known as caste polyphenism that is characteristic of many social insects [70]. Differences in gene expression and alternative splicing between castes has been a topic of intense research interest because it provides a striking example of environmentally-induced phenotypic plasticity. Caste-differential expression has been widely investigated in advanced eusocial honey bees [12, 13, 14] and ants [15, 16]. More recently, high-throughput RNA sequencing technology (RNA-Seq) has been applied to profile expression in species representing a wider array of insect sociality, including an incipiently social small carpenter bee, an intermediately social bumble bee [17] and two primitively eusocial species of *Polistes* wasps [59, 18]. New RNA-Seq data described in this study represent transcriptome data for a third *Polistes* species, facilitating the discovery not only of caste-differentially expressed genes in *P. dominula*, but also of conserved *Polistes*-specific genes.

We performed two lanes of Illumina paired-end RNA-Seq on mRNA isolated from heads of individual adult workers and active egg-laying queens (six replicates per group, from the same population as the male used for genomic DNA sequencing). The RNA-Seq reads were then mapped to 17,888 iLoci using Bowtie [60], with most libraries mapping at an efficiency of 80%, after which iLocus abundances were estimated using RSEM [61] and tested for differential expression using EBSeq [62] [62] (methods and quality control described in detail in **SI Methods**). We identified 381 iLoci differentially expressed between queens and workers (**Figure 2.2A**), 100 lacking annotated gene models, 276 containing a single annotated gene, and 5 containing multiple genes. The majority of the 381 differentially expressed iLoci (231; 60%) are up-regulated in workers. Other

reports that also focused on head or brain gene expression from *Polistes* [59, 18] and honey bees [13] also found the majority of genes are worker-biased in expression. The skew towards worker-biased expression could reflect differences in behavioral flexibility and/or cognitive demands of workers compared to egg-laying queens [71].

Differentially expressed iLoci are significantly enriched for Gene Ontology functions in fatty acid metabolism, neurotransmitter activity, and amino acid metabolism when compared to the background set of all *P. dominula* gene models (**Figure 2.2B**). Previous studies on caste-related gene expression in other *Polistes* species have also identified consistent differences in the expression of genes related to lipid metabolism [59, 72, 48]. These data contribute to a growing base of information suggesting the expression of deeply conserved genes (i.e. a "genetic toolkit") related to metabolism is related to caste differences and may play a key role in the evolution of caste-containing insect societies [73].

All previously examined insects show evidence of large amounts of alternative splicing, including other social insect genomes [74, 19]. As expected, we found evidence for alternative splicing in 1,743 of the *P. dominula* gene models. In particular, *via* transcript mapping and scanning for the two major types of alternative splicing (intron retention and exon skipping, see **SI Methods**) we uncovered 1,616 intron retention events in 1,135 genes and 1,720 exon skipping events in 884 genes (see **SI Results**). 859 genes show only intron retention, 608 genes show only exon skipping, and 276 genes show both. However, analysis with Cufflinks and Cuffdiff [75] reported no cases of caste differential splicing, suggesting alternative splicing is not related to adult caste differences, at least in heads, of *P. dominula*.

Recently, there has been growing interest in the potential for "novel", or taxonomically restricted genes in the evolution of novel phenotypes and in particular, eusociality [21]. We also used our transcriptome data, in conjunction with previously published data for other *Polistes* species, to search for well-supported *Polistes*-specific genes. Our data

represent the third published transcriptome dataset for a *Polistes* species, together with the transcriptomes of two New World species, *Polistes metricus* [59] and the Neotropical Polistes canadensis [18]. In the Polistes canadensis study, the authors identified a large number of novel transcripts (approximately 50% of sequenced transcripts) with no similarity to any known sequence and suggest that novel genes may be related to the evolution of caste differences in social insects [21]. We performed a more in-depth exploration of the three transcriptomes in order to identify *Polistes*-specific transcripts that were shared by all three species. Such transcripts are much more likely to represent true protein-coding genes because they are conserved across species and there is evidence of their expression in multiple species. Considering P. dominula transcripts with an open reading frame of at least 80 aa, we found 19,173 transcripts with no significant similarity to Hexapoda sequences. Only 144 of these transcripts have translation products that are conserved between all three *Polistes* transcriptomes. Of the 144 conserved shared transcripts, 118 are found in the annotated genome assembly, aligning to 93 different iLoci (Figure 2.2C). Only 10 of these 93 iLoci also have evidence of *Polistes*-specific genes from the genome annotation (in the form of gene models without matches to protein databases), and even in these 10 cases there is little agreement between transcript alignment structure and predicted gene structure (Table S11). These results suggest that while single lines of evidence may offer hints of clade-specific genes, very few cases are well supported when subjected to multiple lines of inquiry. This confirms a recent study in ants that uncovered evidence for very few shared, genus-specific genes and more unique species-specific gene, some of which are likely bioinformatics artifacts [16].

There are conflicting reports on the association of novel transcripts in caste differences in *Polistes*. Transcriptomic comparisons from *P. canadensis* adults suggested novel transcripts are more likely to be caste-biased in expression [18], whereas novel transcripts from *P. metricus* larvae did not show this caste-bias [59]. In the current study, we found that 77 out of 93 iLoci (83%) associated with the 144 well-supported *Polistes*-specific

transcripts are caste differentially expressed (significantly overrepresented, Fisher's Exact Test p < 2.2e-16), 34 of which (44%) are up-regulated in workers. In addition, eight of the 10 iLoci containing both unmatched transcripts and unmatched gene models are caste differentially expressed, with 4/8 up-regulated in workers. These results are consistent with data from P. canadensis [18] suggesting novel genes are more likely to be caste-biased in expression in adults. The fact that a similar relationship between caste-biased expression and novel genes was not found in P. metricus larvae suggests there could be different functions for novel genes across species and/or life stages.

DNA methylation system is greatly reduced in *P. dominula*.

There has been great interest in the role of epigenetics in eusociality, and data from honey bees has generated considerable interest in the potential role of DNA methylation in the regulation of gene expression during the development of queen and worker castes [20]. We used the aforementioned genome and transcriptome data from *P. dominula*, along with newly generated whole genome bisulfite sequencing (methylome) data, to probe the presence and extent of caste-association of DNA methylation in the independently evolved social paper wasps.

A full complement of DNA methyltransferases, Dnmt1, 2, and 3, is considered to be necessary for a fully functional DNA methylation system [20]. Dnmt1 is typically considered as the "maintenance" methyltransferase involved in maintaining consistent methylation across cell divisions and generations [20]. Dnmt2 is thought to be involved mainly in the methylation of transfer RNAs. Dnmt3 is the "de novo" methyltransferase, and has been suggested to be related more to environmentally-responsive DNA methylation that occurs within the lifetime of an individual [20]. Other canonical methylation-related proteins include MBD (Methyl-CpG-binding domain protein) and the demethylation enzyme TET (Ten-eleven translocation methylcytosine dioxygenase) [20]. We used BLAST to identify sets of homologs for each of these genes and subjected these sets of sequences

to molecular phylogeny analysis to determine copy numbers of each of these five major DNA methylation related genes.

All previously sequenced Hymenoptera possess a full complement of DNA methyltransferases [76], except the recently sequenced *Polistes canadensis* which lacks *Dnmt3* [42]. Our MAKER annotation workflow (augmented by manual annotations as well as low stringency similarity searches for potentially incomplete or highly diverged homologs) also uncovered no Dnmt3 gene, but did identify one Dnmt1 gene (as in ants [35]), as opposed to two in honey bees [77] and one Dnmt2 gene, as well as genes encoding MBD and TET homologs (summarized in Figure 2.3A). To further investigate whether the absence of a Dnmt3 gene model might represent a gene loss, we examined available Hymenoptera genomes for shared synteny in the region harboring *Dnmt3* in *Apis mel*lifera. Results show that the Dnmt3 locus is within a syntenic block encompassing at least an additional two genes upstream and two genes downstream, conserved in bee and ant genomes. Synteny analyses were conducted with the SynFind and associated tools within the CoGe platform (https://genomevolution.org/coge/; [78]). Sample genome alignments are shown in **Figure 2.3B**. The first upstream and the two downstream genes co-localize in a 90kb region on scaffold0086 of our P. dominula assembly, preserving the syntenic block (the leftmost gene of the bee-ant syntenic block is preserved on scaffold0049 but would be at least 235 kb away if this scaffold were to align upstream of scaffold0086). However, there is a conspicuous absence of any similarity to Dnmt3 in the syntenic region of the P. dominula genome (Figure 2.3B). Intriguingly, across the remaining Hymenoptera species, the region upstream of the annotated $Dnmt\beta$ genes encoding the conserved C-terminus of the methyltransferase is highly variable in size and gene structure annotation is unclear (including annotation of the possibly overlapping upstream gene). The Nasonia vitripennis Dnmt3 protein is highly diverged at the Nterminus, and although the other genes of the bee-ant syntenic block are highly conserved in Nasonia, they are widely spread in the genome. These results suggest that the Dnmt3 locus, and by extension, perhaps some functional aspects of DNA methylation systems in general, are not highly conserved in different lineages of Hymenoptera.

In addition to the lack of genome sequence evidence for a functional Dnmt3 gene in $Polistes\ dominula$, we found no significant similarity between Hymenoptera Dnmt3 sequences and transcripts from any of the three Polistes species' transcriptomes [59, 18] (tblastn search with -evalue 1e-8). The lack of any Dnmt3 transcripts in the three congeners strongly suggests this gene has indeed been lost across the genus Polistes. Further work on additional species will be necessary to determine whether this loss is common to the entire paper wasp family Vespidae.

Along with the loss of Dnmt3, whole genome bisulfite sequencing of one pool of queen and one pool of worker heads revealed a dramatic reduction in DNA methylation in P. dominula compared to other Hymenoptera. This is in contrast to previous reports suggesting the presence of typical amounts of DNA methylation in P. dominula, but these reports used a less reliable method for estimating DNA methylation based on a methylation-sensitive restriction enzyme assay [51, 52]. Through a complete and uniform reanalysis of previously published Hymenoptera bisulfite sequencing data (from honey bees, two ant species, *Polistes canadensis*, and *Nasonia vitripennis*) we were able to make a reliable comparison of DNA methylation levels in P. dominula to those of other Hymenoptera (described in **SI Methods**). Overall, levels of DNA methylation in P. dominula are more than two orders of magnitude lower than in other Hymenoptera. This includes the congeneric *Polistes canadensis*, which, although showing lower levels of methylation than ants and bees, still has two orders of magnitude more methylated CpG sites than P. dominula (Figure 2.3C, SI Results). We uncovered only 124 and 158 CpG methylated sites, respectively in the queen and worker samples; this is in stark contrast to tens of thousands of sites uncovered in all of the other Hymenoptera species (Figure **2.3C**). Similar to other insects [79], most methylated sites were found within genes (74) and 89 sites in queen and worker samples, respectively); see SI Results. Strikingly,

methylation was targeted to the same seven genes in both queen and worker samples (**Figure 2.3D**), and several of these have putative functions related to DNA binding. Thus, there were zero caste differentially methylated genes, and great similarity between castes, even at the level of which cytosines within the seven genes were methylated. Of the 101 total methylated cytosines within the seven genes, 62 (61%) of the same methylated cytosines were shared between both castes (Figure 2.3D). This result is again in contrast to studies from both bees [53] and ants [64], which reported hundreds of caste differentially methylated genes between queen and worker castes. The fact that nearly identical methylation patterns were found in just a few genes, but consistently across castes, suggests the extremely low level of DNA methylation we describe in P. dominula is real and may be of some functional significance. We suggest that, despite a massive reduction in de novo methylation in paper wasps, there may have been selection to retain "maintenance methylation", likely via the action of Dnmt1, for a few key genes. This idea is supported by the observation that five out of the seven P. dominula methylated genes also showed strong methylation in P. canadensis (SI Results), and homologs of three of these seven genes in Apis mellifera are methylated consistently across multiple independently published experiments.

Our methylome data from *P. dominula* also suggest no clear connection to dynamic gene expression patterns: none of the seven methylated genes is differentially expressed between castes; two out of the seven show some evidence of alternative splicing, but not caste differential splicing (PdomGENEr1.2-09385 and PdomGENEr1.2-09184). Although *P. canadensis* also shows some evidence of a reduced methylation system (loss of *Dnmt3* and fewer methylated CpG sites and methylated genes than bees and ants [42], the reduction in *P. dominula* is much more striking. This suggests reduced DNA methylation systems may be a general characteristic of paper wasps, but that there has been even further reduction of these systems in the *P. dominula* lineage relative to some of its congeners.

These data raise intriguing questions about the importance and function of DNA methylation in insects. DNA methylation systems have also been dramatically reduced in other insect lineages (e.g. *Drosophila* flies and *Tribolium* beetles), the shared feature being a loss of *Dnmt3* and large reduction in overall levels of DNA methylation [80]. Furthermore, there are other insects where *Dnmt3* is not present, but moderate levels of DNA methylation remain [81, 42]. Thus, DNA methylation is not clearly related to gene regulation in some insects [82] and even some social insects, suggesting other types of epigenetic mechanisms such as histone modifications [83] or microRNAs may be more important. Our data also highlight the surprising lability of epigenetic mechanisms even within an insect lineage (Hymenoptera) and do not support the idea that phenotypic plasticity afforded by DNA methylation is required for the evolution of castes in social insects [55].

We also examined patterns of occurrence of CpG dinucleotides in the *P. dominula* genome, because segmental ratios of observed to expected (o/e) CpG frequency have been used as an indicator of regional DNA methylation status, based on the assumption that highly methylated regions are characterized by mutational loss of methylated cytosines [84]. The distribution of CpG o/e in *P. dominula* is similarly broad as that of other Hymenoptera, but lacking the bimodal distribution characteristic of the measure in bee coding regions (**Figure S8**). Use of this measure as an indicator of methylation status in *P. dominula* would have incorrectly inferred the presence of numerous methylated genes. Thus, the CpG o/e measure does not accurately reflect the true methylation status of the *P. dominula* genome based on bisulfite sequencing, a much more direct and sensitive method for detecting actual site-specific methylation. It is conceivable that CpG depletion is still correlated with historical (not modern) patterns of DNA methylation, and this is reflected in the fairly typical CpG o/e distribution in *P. dominula* (**Figure S8**). Because appreciable levels of DNA methylation are found in a wide variety of other Hymenoptera (**Figure 2.3A**), it is likely that reduced DNA methylation is a derived

condition in *Polistes*, but more data on additional species are needed to understand when and why reduced DNA methylation evolved in vespid wasps.

Aculeate Phylogeny.

The genome of *Polistes dominula* provides a beneficial complement to the genomes of several species of aculeate Hymenoptera already published. Together, these genomes are a powerful comparative genomics resource for identifying what is conserved and what is unique among the primary aculeate lineages. A delineation of the phylogenetic relationships between these lineages is a fundamental component for analysis and interpretation of evolved traits, and yet consensus regarding the phylogeny of Aculeata remains elusive [85, ?]. A study using molecular data from 4 loci in 64 taxa placed bees (superfamily Apoidea) as sister to scoliid and bradynobaenid wasps [86], while a more recent study involving analysis of 308 genes from 19 taxa found ants (family Formicidae) to be sister to bees [85].

Because our current work describes the one of the first published complete genomes of a vespid wasp, we sought to use these data to investigate the phylogenetic grouping of *Polistes* proteins relative to orthologs in bees, ants, and the non-aculeate wasp *Nasonia* as an outgroup. Using conserved single-copy orthologs present in *Apis mellifera* (bee), *Harpegnathos saltator* (ant), *Polistes dominula* (paper wasp), and *Nasonia vitripennis* (non-aculeate outgroup), we inferred a phylogenetic tree for each gene using these four representative protein sequences (see **SI Methods**). We observed all possible topologies in the 2,077 gene trees: bees and ants as closest neighbors in 889 trees (43%), *Polistes* and bees as closest in 696 trees (34%), and *Polistes* and ants as closest in 492 trees (24%). Although the most common topology (bees and ants as closest) agrees with the results of the most recent, transcriptome-based phylogenetic analysis of aculeates (45), there was definitely not a clear consensus from our data. Based on our analysis, the protein-coding genomes of the published Hymenoptera do not yet provide a definitive

answer to the question of the phylogenetic relationship of bees, ants, and vespid wasps. Additional aculeate genomes, including more representatives of the Vespidae [43] and other wasp families, may help to better resolve aculeate relationships in the future.

Conclusions

This paper provides valuable and comprehensive genomic resources for one of the major lineages of eusocial insects, the vespid wasps, represented by the behavioral model species Polistes dominula. The P. dominula genome is a relatively compact (250Mb) genome with little repetitive DNA, as well as low GC content, in comparison to other Hymenoptera. Transcriptomic analyses revealed several hundred genes with caste-related expression, with functions related to fatty acid and amino acid metabolism and neurotransmitter activity. In addition, we identified several *Polistes*-specific genes, several of which also show differential expression between queen and worker castes. Together, these data provide some support for the roles of both conserved genes and novel genes in the evolution and maintenance of caste differences in social wasps [21, 73]. The most surprising finding from our P. dominula -omics data was clear evidence of a striking reduction in the DNA methylation system. P. dominula have a reduced complement of DNA methylation enzymes, including a loss of the de novo methyltransferase Dnmt3, as well as extremely reduced levels of DNA methylation in the genome—with evidence for just over 100 methylated sites in only seven genes. In addition, there was no relationship between DNA methylation and caste-related gene expression, methylation, nor alternative splicing. There has been great interest and research activity related to the potential role of DNA methylation in the regulation of caste differences and caste evolution in eusocial insects [54, 55]. Our data are novel in that they suggest P. dominula possesses the most reduced DNA methylation system known for any eusocial insect, but there are other examples of non-social insects with similarly reduced methylation systems, including *Drosophila*. These data add to growing evidence for a surprising amount of lability of epigenetic mechanisms in insects, and suggest DNA methylation *per se* is not generally related to the evolution of castes in social insects. These genomic, transcriptomic, and epigenomic data on a primitively eusocial vespid wasp open up exciting new possibilities for comparative genomics of social evolution. Comparisons both across eusocial lineages and within lineages have the potential to provide new insights into the roles of conserved genes and pathways, novel genes, and epigenetic mechanisms in social evolution [87].

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Data Accessibility

Raw Illumina sequences are available from the NCBI Short Read Archive under the following accessions: accession SAMN02584905 for whole genome shotgun reads; accessions SAMN03940809-SAMN03940820 for RNA sequence reads; and accessions SAMN03946123 and SAMN03946134 for bisulfite sequence reads. The genome assembly is available from GenBank under the accession GCA_001465965.1, and the transcriptome shotgun assembly is available from GenBank under the accession GEDB000000000.1.

Additional supporting data, analysis documentation, and supporting code have been deposited in the figshare archive and in several GitHub repositories, as described at https://pdomgenomeproject.github.io/.

Sequences, annotations, and alignments are also available through PdomGDB, an integrated data resource including a genome browser, a BLAST server, and a community annotation portal (see http://goblinx.soic.indiana.edu/PdomGDB).

Author Contributions

All computational analyses of the data were done by DS and VB at Indiana University. AB assisted with the assembly of the *P. dominula* transcriptome and with differential expression analysis. AS contributed to preliminary identification of transposable and repetitive elements, and to analysis of telomere-related genes and sequence motifs. KG contributed template Perl code for preliminary analyses of CpG o/e. AT conceived of and oversaw the project, interpreted results, and collected and processed wasp samples. DS, VB, and AT wrote the manuscript.

Figures

Figure 2.1

A. Best supported molecular phylogeny of the eusocial aculeate Hymenoptera based on recent transcriptome studies and analysis of 2,077 conserved genes reported in this study. B. Stacked bar plot showing genome content of *Polistes dominula* broken down by the proportion occupied by various categories of gene content and conservation, compared to another paper wasp (green labels), two bees (black labels), two ants (red labels), and the outgroup *Nasonia vitripennis*. See also Figure S5. C. Stacked rug plot showing nucleotide composition of long genomic sequences from 3 bees (in black), 2 ants (in red), the outgroup *Nasonia vitripennis* (in blue), and 2 paper wasps (in green). Each vertical bar represents a chromosome, linkage group, or scaffold at least 1 Mb in length. Photo credits: *N. vitripennis* by E. Cash and J. Gibson; *P. dominula* by S. McCann; *S. invicta* and *A. mellifera* by A. Wild.

Figure 2.2

A. Heatmap of expression values of 367 differentially expressed interval loci (iLoci). The blue color indicates overexpression, while the yellow indicates underexpression. 212 iLoci (58%) of the differentially expressed iLoci are overexpressed in workers. B. Bar chart showing the representation of eight GO functional categories as a proportion of differentially expressed iLoci (blue bars) versus all iLoci (red bars), determined by an enrichment analysis to be overrepresented in differentially expressed iLoci. C. Putative *Polistes*-specific genes, defined as unmatched transcripts with significant pairwise protein-level similarity among three Polistes species (Pd for *P. dominula*, Pc for *P. canadensis*, and Pm for *P. metricus*). Because of variation in gene copy number and number of alternative transcript isoforms, the number of transcripts in each intersection of the diagram is different for each species, and only the smallest number is shown. For example, the

3-species intersection consists of 144 transcripts from *P. dominula*, 136 transcripts from *P. canadensis*, and 95 transcripts from *P. metricus*.

Figure 2.3

A. Copy number of five methylation-related genes in the primary Hymenoptera lineages. B. Evidence for shared synteny around the Dnmt3 locus in bee, paper wasp, and ant genomes. Colored bars represent conserved coding regions between *Polistes domin*ula (top track) and Apis mellifera (black blocks), Bombus terrestris (grey blocks), and Camponotus floridanus (red blocks). Regions of similarity are largely collinear (shown for the Apis mellifera to Camponotus floridanus comparison by the blue lines connecting the similar blocks). Gene models are shown by arrow structures with coding exons in green, UTRs in blue, and introns as thin lines. Each gene is denoted by a numbered box as follows: 1) GenBank protein entries XP_006568814 (26S proteasome non-ATP regulatory subunit 6-like), 2) XP_006568813 (uncharacterized), XP_00658716 (Dnmt3), 4) XP_006568806 (polynucleotide 5'-hydroxyl-kinase NOL9-like), and 5) XP_0065688112 (histone lysine demethylase PHF8-like) in A. mellifera (linkage group LG2, GenBank $NC_{-007071}$), from left to right. The lack of similarity blocks around 50K on the P. dominula scale demonstrates the postulated loss of Dnmt3 in this species. C. Bar chart showing the number of CpGs with a high level of support for methylation from bisulfite sequence data. The number of highly supported methylation sites for each species is based on pooled reads for all available samples, and is shown for one bee (black), two ants (red, note that "worker" refers to "minor workers" in C. floridanus), two paper wasps (green), and N. vitripennis (blue). D. Gene models for the 7 P. dominula methylated genes (with corresponding Gene ID numbers and putative annotations based on best BLAST hits). Approximate locations of each of the 124 highly supported methylation sites within each gene are indicated with a line-dot symbol, with sites methylated in both queen and worker samples (n=76) indicated in black, sites methylated in just the

queen sample in green (n=18), and sites methylated in just the worker sample (n=30) in orange.



Figure 2.1



Figure 2.2



Figure 2.3

CHAPTER 3. PARSEVAL: PARALLEL COMPARISON AND ANALYSIS OF GENE STRUCTURE ANNOTATIONS

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Abstract

Background: Accurate gene structure annotation is a fundamental but somewhat elusive goal of genome projects, as witnessed by the fact that (model) genomes typically undergo several cycles of re-annotation. In many cases, it is not only different versions of annotations that need to be compared but also different sources of annotation of the same genome, derived from distinct gene prediction workflows. Such comparisons are of interest to annotation providers, prediction software developers, and end-users, who all need to assess what is common and what is different among distinct annotation sources. We developed ParsEval, a software application for pairwise comparison of sets of gene structure annotations. ParsEval calculates several statistics that highlight the similarities and differences between the two sets of annotations provided. These statistics are presented in an aggregate summary report, with additional details provided as individual reports specific to non-overlapping, gene-model-centric genomic loci. Genome browser styled graphics embedded in these reports help visualize the genomic context of the annotations. Output from ParsEval is both easily read and parsed, enabling systematic

identification of problematic gene models for subsequent focused analysis.

Results: ParsEval is capable of analyzing annotations for large eukaryotic genomes on typical desktop or laptop hardware. In comparison to existing methods, ParsEval exhibits a considerable performance improvement, both in terms of runtime and memory consumption. Reports from ParsEval can provide relevant biological insights into the gene structure annotations being compared.

Conclusions: Implemented in C, ParseEval provides the quickest and most feature-rich solution for genome annotation comparison to date. The source code is freely available (under an ISC license) at http://parseval.sourceforge.net/.

Background

It was only a decade ago when annotating a eukaryotic genome required years of extensive collaboration and millions of dollars of investment. Since then, the tremendous rate at which the cost of DNA sequencing has been dropping as well as increased accessibility to gene prediction software are placing genome sequencing and annotation well within the reach of most single investigator biology laboratories. As a result, proliferation of distinct annotation sets corresponding to the same genomic sequences is becoming increasingly common. Annotation sets for a particular genome can accumulate in a variety of scenarios. When developing gene prediction software, it is common to test the software on a genomic region for which a high-quality reference is available, running and re-running the software and comparing the resulting predictions against the reference. Community groups providing annotation for species- or clade-specific genomes typically release updated annotations following the initial release. Affordable transcriptome sequencing provides individual labs with data to specifically improve annotations for particular genes of interest, for example with respect to alternative splicing. In each of these scenarios, multiple annotations associated with a common set of genomic sequences

Table 3.1 Annotation Comparison Methods

Method	Pros	Cons
Manual comparison	minimal overhead	extremely tedious; error prone; unscalable
Genome browser	intuitive interface; visual assessment of indi-	visual assessments imprecise; extensive over-
	vidual loci	head; little or no automation
Eval	detailed statistics; visual assessment of statis-	older software; relatively slow; only summary
	tic distributions; scales fairly well for large	statistics are reported, while stats for individ-
	data sets; can compare multiple predictions to	ual loci are discarded
	a single reference	
ParsEval	detailed statistics provided, not only as a sum-	only capable of comparing a single pair of an-
	mary but for individual loci as well; scales well	notations
	for large data sets; fast, efficient, and portable	

Various approaches to comparing alternative sources of gene structure annotations, with a brief description of the associated pros and cons.

require comparative assessment.

A variety of comparison methods exist, but none can fully address the growing needs of the community (see Table 3.1). Manual comparison approaches can trivially be ruled out as slow, tedious, error prone, and hopelessly unscalable. Although genome browsers have had a huge impact by making gene annotations accessible to a wide variety of scientists, they likewise do little to provide the automation and precision needed in whole-genome annotation comparisons. Large genome sequencing projects and centers have certainly developed in-house scripts and pipelines over the years to address this need. However, these pipelines are typically not standardized, not openly shared, and do not migrate well.

Tools such as the Eval package [23] and the GFPE program [24] represent some of the earliest efforts to provide a reusable, easy-to-use annotation comparison tool to the community. Eval in particular stands out based on the amount of detail provided by its reported comparison statistics and by the ability to visualize the distributions of these statistics. Eval takes as input annotation files in Gene Transfer Format (GTF) and calculates a rich set of descriptive statistics summarizing the differences between the annotations. Because whole-genome annotations typically include thousands (or tens of thousands) of genes, these statistics are intended to condense the information into a comprehensive yet concise summary (at the resolution of entire sequences or sets of sequences), facilitating targeted improvement of gene prediction software. Unfortunately,

this condensing process discards large amounts of valuable information at the resolution of individual gene loci, making the tool unsuitable for analyses that target a particular gene, sets of genes, or gene loci with characteristics of interest from within a larger set of genes. Such locus-resolution comparisons are useful not only to software developers and annotation producers who need to know whether their software has distinct advantages or disadvantages, e.g., favoring long over shorter gene models on average, or failing in untranslated region (UTR) prediction, but they are of primary interest for specialists concerned with a particular gene family or pathway.

Motivated by a need for genome-scale evaluations with locus-scale detail, we developed ParsEval, a program for comparing and analyzing distinct sets of gene structure annotations for the same input sequences. The program is designed to incorporate all of the benefits of existing methods while addressing their shortcomings. ParsEval identifies differences in exon/intron assignments and in coding sequence (CDS) and UTR designations, at both feature-level (exon, CDS segment, UTR segment) and nucleotidelevel resolution. The output consists of a set of commonly used statistics that provide quantitative measures of agreement when comparing predicted gene structures against a standard reference [88, 89, 6]. This output is presented in a detailed report for each gene locus, supplemented with genome browser styled graphics to enable additional visual assessment and analysis of the annotations. The statistics are also presented in a single summary report that aggregates the statistics across all loci, providing a condensed high-level view of the similarity between the two sets of annotations. For gene loci that include alternatively spliced genes or overlapping genes (or both), ParsEval determines the optimal matching of reference transcripts to prediction transcripts, and additionally reports any novel transcript predictions that have been identified.

Implementation

Overview

ParsEval is a gene annotation comparison and analysis tool, designed with a focus on speed, resource efficiency, and portability. The program takes as input a pair of gene structure annotations corresponding to the same sequence (in GFF3 format [90]), analogous to two separate annotation tracks one might see in a genome browser. For comparison purposes, the first set of annotations is treated as the *reference* while the other is treated as the *prediction*, although ParsEval makes no assumptions regarding the respective quality of the two annotation sets. The output of the program is a set of reports containing common comparison statistics intended to highlight relevant similarities and differences between the two sources of annotation.

ParsEval first loads the annotation data into memory, identifies start and end coordinates for gene loci, and associates each gene annotation with a single locus. Next, the program does a comparative assessment of the gene annotations for each locus, calculating and storing a variety of informative similarity statistics. Finally, ParsEval generates reports providing a detailed readout of these statistics.

Implemented in ANSI C, ParsEval is fast, memory efficient, and portable, designed to run on all POSIX-compliant UNIX systems (Linux, Mac OS X, Cygwin, Solaris, etc.). Most of the analysis code is implemented with shared memory parallelization, providing additional performance gains when running on multicore processors that are becoming increasingly common in commodity hardware. ParsEval's only external dependency is the GenomeTools library [91], which provides an API for generating annotation graphics with AnnotationSketch [92], as well as implementations of a variety of data parsers and dynamic data structures.

Gene locus identification

Comparative analysis of two sets of gene annotations requires determining how annotations from one set correspond to annotations from the other, as well as the genomic coordinates (the *gene locus*) that should be considered in each comparison. For rare cases in which a single reference annotation and a single prediction annotation line up perfectly, determining the gene locus and the corresponding genes is trivial. However, in most cases this task is complicated by a variety of factors. For example, a single gene prediction workflow may annotate multiple genes at a single location, so one must determine how to associate these annotations with corresponding annotations from an alternative source. Furthermore, when one or more gene annotations from one source overlap with multiple annotations from another source, one must determine how to compare these gene annotations and which coordinates to include in the comparison.

One common approach involves designating one set of annotations as the reference set and then using the coordinates of each reference gene annotation to define a distinct gene locus to serve as the basis for subsequent comparison (see Figure 3.1). However, this approach is unfavorable for several related reasons. First, reference gene annotations that overlap are handled separately, when it makes more sense to associate them with the same locus and handle them together. Second, it forces a quality judgment between the two sets of annotations when their relative quality is often unknown. The two sets of annotations likely include complementary information, and unless there is a clear distinction in quality between the two, choosing one as a reference discards clearly related information from the other. Third, relevant information from predicted gene models that extend beyond the boundaries of the corresponding reference annotation is ignored.

Although ParsEval uses the terms reference and prediction to distinguish between the two sets of annotations, both are considered equally when identifying gene loci. Each gene annotation corresponds to a node in an interval graph G. There is an edge between two nodes G_i and G_j if the corresponding gene annotations overlap (see Figure 3.2).

Each connected component in G then corresponds to a distinct gene locus, which we define as the smallest genomic region containing every gene annotation associated with the corresponding subgraph. Defining a gene locus in this way makes no assumptions as to the relative quality of the two sets of annotations, and ensures that no potentially relevant data are discarded. Furthermore, according to this definition each gene locus is independent, enabling the subsequent comparative analysis tasks to run in parallel.

Gene structure representation

To facilitate analysis at each gene locus, ParsEval converts GFF3 annotations for each gene into a character string representing the annotated gene structure (a model vector). This model vector is similar to a sequence in Fasta format, except instead of using the alphabet $\{A, C, G, T\}$ to represent chemical composition at each nucleotide, the alphabet $\{C, F, G, I, T\}$ representing gene structure is used: C for coding sequence, F for 5'-UTR, T for 3'-UTR, T for introns, and T for intergenic sequence. Using this alphabet, each transcript can be represented by a single model vector. ParsEval uses these model vectors when comparing reference and prediction gene annotations.

In many cases, a single pair of model vectors (one for the reference, one for the prediction) is sufficient to fully represent annotated gene structure at a given locus. This is certainly true when both the reference and the prediction annotate a single gene with a single mRNA product at the locus. But even if the reference (or the prediction) annotates multiple genes or transcripts, non-overlapping annotations can be encoded in the same model vector and compared simultaneously with corresponding annotations from the other data set. However, if either the reference or the prediction contains annotations for overlapping transcripts, either because of alternative splicing or because of overlapping gene models, a single pair of model vectors is insufficient to represent the complete annotated gene structure at that locus. In these more complicated cases, the reference or the prediction or both will be associated with multiple model vectors.

Thus, the algorithmic requirement is to represent all annotated transcript structures in the locus using the smallest number of model vectors.

This problem reduces to a common problem in graph theory known as the maximal clique enumeration problem [93]. We treat each transcript as a node in an undirected graph and place an edge between two nodes if the corresponding transcripts do not overlap (unlike the locus identification step, reference annotations and prediction annotations are handled separately in this step). Each maximal clique (maximal fully-connected subgraph) in this graph corresponds to a set of transcripts that do not overlap and can therefore be collapsed into a single model vector. ParsEval uses the Bron-Kerbosch algorithm [93] to enumerate all maximal transcript cliques, first for the reference and then for the prediction. A model vector is generated for each clique, after which ParsEval compares all reference model vectors with all prediction model vectors.

Comparative analysis of annotations

Given a pair of equal-length model vectors representing a pair of gene structure annotations at a given locus, ParsEval computes a variety of comparison statistics to measure the level of agreement between the pair of annotations. Calculated at different levels of resolution, these statistics provide a detailed assessment of similarity between the reference and the prediction. At the resolution of distinct annotation features, ParsEval calculates the sensitivity and specificity as described in [88], the F1 score as described in [89], and the annotation edit distance as described in [6, 94]. These statistics are calculated for exons, CDS segments, and UTR segments. Note that for a prediction feature to be considered a true positive, ParsEval requires both the start and end coordinates to match the reference perfectly.

At the nucleotide-level resolution, ParsEval also calculates the sensitivity, specificity, F1 score, and annotation edit distance, as well as the simple matching coefficient and the correlation coefficient as described in [88]. These statistics are calculated for coding

nucleotides (CDS) and untranslated exonic nucleotides (UTR). Overall identity at the nucleotide level, of which the simple matching coefficient is a generalization, is also computed.

For complex loci requiring multiple comparisons, the locus report includes an aggregate summary of the similarity statistics at the locus level in addition to the reports for each individual comparison. This locus-level summary also includes the splice complexity statistic [6], which ParsEval computes and reports for both the reference and the prediction at the locus level.

Based on the computed statistics, each comparison is classified in terms of similarity. A comparison is classified as a perfect match if the model vectors (and by implication the annotated gene structures) are identical. A comparison is classified as a CDS structure match if the comparison is not a perfect match, but there is perfect agreement in terms of CDS structure. A comparison is classified as an exon structure match if there are differences in the coding sequence that nevertheless preserve exon structure (as resulting from different start and/or stop codons). A comparison is classified as a UTR structure match if there are differences in CDS and exon structure, but the UTR structures are identical. All other comparisons are classified as non-matches.

Note that, as with feature-level statistics, match classifications require perfect agreement. For instance, a pair of annotations may have very similar CDS structures, and this will be reflected in the nucleotide-level CDS statistics. However, if the CDS structure tures are not precisely identical, the comparison will not be classified as a CDS structure match.

As comparison statistics are computed on a locus-by-locus basis, ParsEval also maintains a running total of all comparison counts (such as true positives and false positives) from which the statistics are computed. When all loci have been considered, each comparison statistic is then recomputed using these running totals to provide an overall assessment of similarity.

Reporting comparison scores

For each gene locus, comparison statistics are calculated for each corresponding pair of reference and prediction model vectors. If multiple comparisons are required at a locus, however, statistics are not reported for each comparison. The comparisons are ranked using the previously described similarity statistics and are reported so as to ensure each transcript (or transcript clique) is considered at most one time. In cases where there is an unequal number of reference and prediction transcripts (or transcript cliques) associated with a particular locus, some will be labeled as novel or unmatched transcripts, and corresponding statistics are not included in ParsEval's reports.

ParsEval presents the comparison statistics in a collection of reports. The first is a single summary report providing the aggregated statistics for a high-level assessment of similarity, as is standard for tools of this kind. Additionally, ParsEval produces a dedicated comparison report for each individual locus. The detail provided by these locus-level reports is extremely valuable, and ParsEval is the only tool of its kind that preserves and reports comparisons at this level. By default, ParsEval generates these reports in an easy-to-parse and easy-to-read text format. However, ParsEval can also generate the reports as hyperlinked HTML files to facilitate browsing and network-based distribution. Furthermore, ParsEval can supplement HTML reports with embedded PNG graphics providing a genome-browser-like view of each locus' genomic context and enabling visual assessment of the annotations.

If more targeted reporting is desired, ParsEval also provides some filtering features. Using a simple optional configuration file, the user can exclude some gene loci from the reports based on a variety of features: locus length, number of genes, number of transcripts, number of transcripts per gene, number of exons, and CDS length. No comparisons are performed for loci that are filtered out, and thus do not contribute to the reported aggregate summary statistics and comparison classifications.

To facilitate integration of comparison reports with popular genome browsers such

as GBrowse [95] and PlantGDB [96], ParsEval can generate an additional output file (in GFF3 format) containing the coordinates of each gene locus. These genome browsers commonly allow users to anonymously create private custom tracks with uploaded data, which provides the quickest mechanism for integration. Once a track is populated with the uploaded locus data, the user can configure the track configuration so that each locus feature in the track is hyperlinked to the corresponding ParsEval report stored, for example, on that user's local machine (see Figure 3.3). Alternatively, if a more permanent and public solution is desired, a user with administrative privileges for the genome browser can follow standard procedures for populating a new track with the GFF3 data, and then configure the track so that locus features are linked to network-accessible ParsEval reports.

Results and Discussion

We present several use cases to demonstrate ParsEval's capabilities, benchmark its performance, and compare its utility relative to existing methods. The input data for these demonstrations were obtained from a variety of public databases with different respective formatting conventions. Accordingly, all data files were processed and converted to a uniform format before analysis. A detailed description of this conversion process, along with all code and commands used, are provided in the Supplemental Data as well as in ParsEval's source code distribution.

Unless otherwise noted, all use cases and benchmarks described herein were run on a fairly modest desktop computer: a Mac Pro with two 2.8 GHx quad-core Intel Xeon processors and 4 GB of RAM. ParsEval's performance for these demonstrations should therefore be fairly representative of the performance one might expect when running on commodity laboratory or personal hardware.

Use case: predictions vs. gold standard

High-quality gene structure annotations derived from a combination of computational and experimental evidence, and possibly improved with expert manual curation, are indispensably used as "gold standards" for measuring the accuracy of a novel gene prediction method or entire new annotation workflows. Identifying differences between the new method's predictions and such gold standard reference can help identify areas in which the novel method provides or needs improvement. Reports from ParsEval are effective for quickly and clearly identifying such differences.

To demonstrate ParsEval in this context, we reproduced a comparison that was originally published to assess the performance of the AUGUSTUS gene prediction program [97]. In the original study, AUGUSTUS was tested on the h178 data set [98], a set of 178 human genomic sequences, each containing a single gene, for which annotations were available from the EMBL database release 50 [99]. Gene predictions from AUGUSTUS were compared the annotations from EMBL, and sensitivity and specificity scores were calculated at the nucleotide level, the exon level, and the gene level.

We obtained the h178 data set (sequences and EMBL r50 annotations) from [100]. We then used the latest version of AUGUSTUS (2.5.5) to generate gene predictions for the 178 sequences. The data files were reformatted and then compared using ParsEval. Running on a desktop computer, ParsEval generated graphical reports in less than a minute. The summary report provided immediate access to a variety of similarity metrics, including those reported in the original assessment. The sensitivity and specificity values reported by ParsEval are comparable to those reported in the original AUGUSTUS manuscript (see Table 3.2). Differences in the comparison metrics can likely be explained by improvements to the AUGUSTUS program since publication, although the exact reason is elusive since the original AUGUSTUS software is no longer accessible.

Table 3.2 Use case: prediction vs. gold standard

Statistic	AUGUSTUS manuscript	
Coding nucleotide sensitivity	0.93	0.94
Coding nucleotide specificity	0.90	0.99
Exon sensitivity	0.80	0.81
Exon specificity	0.81	0.86
Gene sensitivity	0.48	0.43
Gene specificity	0.47	0.46

Sensitivity and specificity scores for AUGUSTUS gene predictions in comparison to corresponding gene annotations from EMBL database release 50. The first column shows scores as reported in the original AUGUSTUS manuscript. The second column shows scores as computed by ParsEval using predictions from the latest version of AUGUSTUS (2.5.5). Summary reports from ParsEval provide immediate access to a wide variety of similarity statistics, including the ones reported in this table. Differences between the scores reported by the AUGUSTUS authors and the ParsEval authors are likely due to subsequent updates of the AUGUSTUS program since its publication.

Use case: two sets of annotations

When working with genome annotations, there is an increasing variety of cases in which no gold standard is available for comparison. For example, gene annotations for many model species are available from a variety of sources (i.e., UCSC versus Ensembl). The respective quality of these different annotation sets is not always clear, but comparison is still a necessary and fundamental task. Another example relates to genome projects that typically offer multiple releases of gene annotations between each major genome assembly release. Although newer releases may offer marginal improvements over the older ones, neither one can truly be considered a high-quality standard reference for comparison. An additional example relates to the increased affordability of genome sequencing and the number of new and exotic species for which genome sequence is available. Gene annotation software is based on complex statistical models containing many parameters, and it is not always initially clear which parameter values to use up front. Therefore, when annotating a newly sequenced genome, it is common to extract a subset of the genome on which to perform repeated optimization runs to determine the parameter values that should be used subsequently to annotate the entire genome.

In each of these scenarios, multiple annotation sets must be compared, despite hav-

Table 3.3	Use case: two sets of annotations	
Perfect matches	22,333	94.7%
CDS structure matches	0	0.0%
Exon structure matches	0	0.0%
UTR structure matches	83	0.4%
Non-matches	$1{,}174$	5.0%
Total	23,590	100.0%

Results from a ParsEval comparison of gene annotations for *Mus musculus* from two recent releases of the Ensembl database (releases 64 and 65). Release 64 contains 22,507 gene annotations, while release 65 contains 14,486 gene annotations. ParsEval identified 20,362 gene loci using these two data sets, 6,725 of which contained only annotations from release 64. For the 13,637 gene loci for which both release 64 and 65 have annotations, 23,590 comparisons were performed. Each of these comparisons was classified according to how well the annotations from the two releases agreed. This table shows a breakdown of these results.

ing no intuition as to the relative quality of the respective annotations. ParsEval was designed precisely for this type of analysis. Reports from ParsEval provide both an overall summary and locus-level detail, enabling the user to make informed decisions about annotations for individual loci, as well as for annotation sets as a whole.

As a demonstration of ParsEval's capability in this context, we downloaded two recent gene annotation releases (releases 64 and 65) for *Mus musculus* from the Ensemble database [101]. We compared these annotations using ParsEval, which required approximately 3 minutes of runtime on a desktop computer. A brief review of ParsEval's summary report shows that a total of 20,362 gene loci were identified using these annotations (see Table 3.3 for a complete breakdown). Of these gene loci, 6,725 had only annotations from release 64.

23,590 comparisons were performed by ParsEval, of which 22,333 (94.7%) were perfect matches between releases 64 and 65. A small number (83, 0.4%) of comparisons were classified as UTR structure matches. For the remaining 1,174 comparisons (5.0%) that were classified as non-matches, transcripts from release 64 contained an average of 16.47 exons, whereas transcripts from release 65 contained an average of 8.11 exons. A brief review of a handful of selected loci showed that many long transcripts (with many exons) that had been present in release 64 were absent in release 65.

This use case is an ideal demonstration of ParsEval's capabilities. Although the authors have no prior experience working with these particular data sets, a cursory examination ParsEval's reports clearly draw attention to an important fact—between release 64 and 65, changes to Ensembl's annotation pipeline (perhaps different values for parameters that influence joining/splitting annotations, or implementation of stricter filters for gene length) affected approximately 5% of the gene annotations. Not only does ParsEval provide this information in a summarized form, it also provides detailed locus reports enabling users to scrutinize the results on a gene-by-gene basis. This breadth and detail of information is of great benefit to a wide variety of scientists and will empower them to more fully understand the available data and make informed decisions regarding alternative sources of annotation.

Benchmarks

To demonstrate its speed, scalability, and efficiency, we benchmarked ParsEval by analyzing pairs of whole-genome gene structure annotations for four common model organisms representing a wide range of eukaryotic diversity: *Arabidopsis thaliana* (thale cress), *Drosophila melanogaster* (fruit fly), *Glycine max* (soybean), and *Homo sapiens* (human) (see Table 3.4). To give a detailed demonstration of its performance, ParsEval was run 24 times for each species—3 technical replicates while varying the output mode (text and HTML/PNG) and the number of dedicated processors (1, 2, 4, and 8). Reported runtimes were obtained by taking the mean of the 3 corresponding replicates.

Performance in text output mode

ParsEval demonstrated optimal performance when running in text output mode, with runtimes ranging between about 30 seconds to about 4 minutes. Running ParsEval in parallel on multiple processors provided noticeable improvement in runtime for *Drosophila* and human, although no improvement was seen for *Arabidopsis* and soybean.

			TOT	TWOIC O. T DOILDING IND		CC		
	A. t	A. thaliana	D. mei	D. melanogaster	9	G. max		H. sapiens
Reference annotations	L	TAIR9	FlyB	FlyBase 5.39	NCE	NCBI Entrez	UCSC 1	UCSC knownGene (hg19)
Prediction annotations	T/	TAIR10	Ense	Ensembl r65	JGI /	JGI / Phytozome	Н	Ensembl r65
Average runtime (sec)	Text	HTML	Text	HTML	Text	HTML	Text	HTML
n = 1	36.3	859.4	91.1	1,350.5	85.3	1,461.1	294.3	6,422.0
n = 2	32.8	449.2	56.6	859.5	79.4	768.4	181.3	4,089.5
n = 4	30.7	246.5	39.2	633.7	76.5	439.9	130.1	2,751.2
n = 8	29.8	168.7	32.4	546.6	76.3	330.5	108.0	2,323.3
Gene loci	2	24,728	1(10,976	4	47,877		17,865
shared		24,460	1	10,944		37,942		7,779
unique to reference		112		32		3,363		9,569
unique to prediction		156		0		6,572		517
Comparisons	33	31,635	55	22,474	3	38,734		16,168
perfect matches	30,638	96.8%	0	0.0%	2,489	6.4%	2,517	15.6%
CDS structure matches	339	1.1%	0	0.0%	17,450	45.1%	8,269	51.1%
exon structure matches	7	0.0%	22,464	100.0%	26	0.1%	27	0.2%
UTR structure matches	115	0.4%	0	0.0%	647	1.7%	28	0.4%
non-matches	536	1.7%	7	0.0%	18,122	46.8%	5,297	32.8%

perfect gene structure matches, how many were CDS structure matches, and how many were non-matches. All of the results shown in this table were easily obtained from the fly), Glycine max (soybean), and Homo sapiens (human). For each organism, we used ParsEval to compare the two corresponding sets of annotations. Runtimes are shown for both text and HTML/PNG output modes, using 1, 2, 4, and 8 processors. For each organism, we also show the number of gene loci identified, how many were shared between the two sets of annotations, and how many are unique to one set. Finally, we show the number of reported comparisons for each organism and how many were As a demonstration of ParsEval's speed and scalability, we obtained pairs of whole-genome annotations for Arabidopsis thalia summary reports generated by ParsEval.

It is likely that for loci with relatively small and simple gene structures, ParsEval's runtime is bound more by serial I/O related tasks than by actual analytical computations, which would explain why no improvement was observed for the plant species.

Performance in HTML output mode with PNG graphics

Running ParsEval in HTML/PNG output mode increased the runtimes by an order of magnitude, although parallel processing kept these runtimes within a reasonable range (about a half hour for the most intensive comparison) with observed speedup factors ranging from 3 to 5 when using all 8 processors. Because these improvements in runtime were observed for all species, it is likely that ParsEval's runtime is bound primarily by computationally intensive graphics generation tasks when running in HTML/PNG output mode.

Notes on benchmark results

The results of the A. thaliana benchmark were not surprising. Perfect matches and CDS matches account for 97.5% of the comparisons, which makes sense considering that TAIR10 represents minor cumulative updates to TAIR9 (in contrast, perfect matches and CDS matches account for only 4.2% of comparisons between TAIR6 and TAIR10). There were even fewer differences between FlyBase and Ensembl annotations for the D. melanogaster benchmark ($\approx 0.1\%$ of loci), suggesting perhaps that these differences may be the consequence of technical artifacts in one data set or the other.

The results of the other two benchmarks, for *G. max* and *H. sapiens*, were somewhat surprising. In each case, approximately 10% of the comparisons reflected perfect matches between the two annotations (6.4% for soybean and 15.3% for human), while approximately 50% of the comparisons reflected CDS matches (45.1% for soybean and 54.9% for human). Therefore, for the remaining approximate 30% of human genes and 50% of soybean genes, the annotated coding sequence (and the associated polypeptide) is

different depending on the data source. These differences are likely the result of different annotation strategies between the alternative sources of annotation. Regardless, this is an important point of consideration both for consumers and producers of gene structure annotations, and we hope that the ParsEval tool will be a useful asset to a wide variety of scientists that rely on reliable gene annotations for their research.

Performance evaluation in comparison to Eval software

To evaluate ParsEval's performance in comparison to existing methods, we used the Eval tool [23] to repeat one of the previously described use cases. Gene annotations for *Mus musculus* were retrieved from releases 64 and 65 of the Ensembl database, and subsequently analyzed using both Eval and ParsEval. Some small differences were observed in the similarity statistics computed by the two programs, although this was not unexpected as Eval uses a different approach than ParsEval for matching reference annotations to prediction annotations. Also, the two programs provide a different breakdown of the similarity statistics, making a rigorous comparison between the Eval results and the ParsEval results impractical.

Running Eval on the complete data sets exhausted the desktop computer's memory resources after several minutes, so comparison of Eval and ParsEval was only possible after restricting the data sets to annotations for *M. musculus* chromosomes 1 through 10. To analyze these reduced data sets, Eval required an average of 12 minutes 13 seconds and consumed all available memory. On the other hand, ParsEval, running on a single processor, required an average of 1 minute 44 seconds, with memory consumption peaking at approximately 0.5 GB. When run on 4 processors, ParsEval's performance margin increased with an average runtime of 47 seconds.

To ensure that Eval's performance was not being severely affected by the desktop's limited system memory, the comparison was also performed in a high-performance computing environment in which memory could not have been a limiting factor. ParsEval

continued to demonstrate superior performance in this environment as well, although by a slightly less drastic margin. The Eval program required an average of 7 minutes 18 seconds of runtime, while ParsEval required an average of 1 minute 19 seconds using a single processor, or 37 seconds using 4 processors.

These tests conclusively demonstrate two important points regarding the performance of ParsEval relative to Eval: not only is ParsEval markedly faster, but its resource efficiency also makes it much better equipped to run whole-genome comparisons on the laptop or desktop computers one might expect to see in the typical biology lab. The initial runtimes reported herein should be fairly representative of what users can expect to observe when running ParsEval on commodity hardware.

Conclusions

The accessibility of genome annotation tools to an increasingly wider variety of scientists will soon be accompanied by an increased demand for supplementary tools to manage and analyze genome annotations. We address this need with ParsEval, a tool for fulfilling a common, fundamental analytical need for which existing software is lacking. ParsEval is a portable, easy-to-install, and efficient program for comparing gene structure annotations, and facilitates a wide variety of downstream comparative analyses. We demonstrate the speed and scalability of ParsEval, even when working with large eukaryotic genomes. Furthermore, we highlight the capability of the detailed comparison statistics in ParsEval reports to highlight relevant biological trends in the data. We anticipate that ParsEval will enable a wide variety of biologists to more fully take advantage of the vast genome annotation data resources accumulating in their individual labs and in the community at large.

Availability and requirements

Source code for ParsEval is available at http://parseval.sourceforge.net under an ISC license. ParsEval is implemented in ANSI C and is designed to run on all POSIX-compliant UNIX systems (Linux, Mac OS X, Cygwin, Solaris, etc.). Aside from a C compiler with OpenMP support (such as GCC 4.2 or higher), ParsEval's only external dependency is the GenomeTools library [91].

Authors contributions

DS designed and implemented the software and drafted the manuscript. VB supervised the project and provided design and feature suggestions. Both authors conceived the project, edited the manuscript, and approved the final version.

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Figures

Figure 3.1

Associating Annotations with Gene Loci. The black bar provides a scale corresponding to a genomic region for which two sets of annotations are available. Reference annotations for gene structure are represented with red glyphs, while prediction annotations are shown with blue glyphs. Arrows indicate the strand of the gene annotation, and different levels of shading correspond to different gene structure features: dark shading for coding sequence, medium shading for UTRs, and light shading for introns. Green brackets denote gene loci as determined by the common practice of using only the genomic coordinates from reference gene annotations. Orange brackets denote gene loci as determined by ParsEval, which takes into account both reference and prediction annotations when selecting distinct loci for comparison.

Figure 3.2

Locus Identification Using a Gene Interval Graph. Red and blue nodes in this interval graph correspond to reference and prediction gene annotations (respectively) as shown in Figure 3.1. Two nodes are connected by an edge if the corresponding gene annotations overlap. Each connected component in the graph represents a distinct gene locus, defined as the smallest genomic region containing every gene annotation associated with the corresponding subgraph. In this example, nodes representing five reference annotations and four prediction annotations are shown. The four connected components in the graph correspond to four gene loci, for which precise genomic coordinates can be determined from the associated genes (shown in orange brackets in Figure 3.1).

Figure 3.3

Integrating ParsEval Reports with a Genome Browser. Screenshot of the Arabidopsis thaliana genome browser at Phytozome (http://phytozome.net/), with a custom anonymous user track populated by ParsEval output. Boxes in this custom track represent loci identified by ParsEval and are color-coded according to the level of agreement between the two sets of annotations compared (dark red and pastel blue glyphs, respectively). This custom track can easily be configured so that features are hyperlinked to ParsEval reports containing detailed comparison statistics.



Figure 3.1



Figure 3.2



Figure 3.3

CHAPTER 4. ILOCI: ROBUST GENOME ANNOTATION AND ANALYSIS FOR PROVISIONAL GENOME ASSEMBLIES

A paper to be submitted to Genome Biology.

Standage DS, Brendel VP

Abstract

Background: The rate at which new draft genome sequences and corresponding annotations are being produced outpaces the scientific community's capacity to refine these drafts into "finished" reference-quality data resources. Scientists must be able to evaluate newly sequenced genomes in the context of previously published data, requiring summaries of genome content that can be quickly computed and meaningfully compared without the support of a large model organism research community. As annotation quality will necessarily vary within and across data sets, the ability to select subsets of only those data that are well supported is critical for distinguishing technical artifacts from biological effects.

Results: We introduce a new framework for genome analyses based on parsing an annotated genome assembly into distinct *interval loci (iLoci)*. We demonstrate that iLoci provide an alternative coordinate system that is robust to changes in assemblies and annotations, and facilitates granular quality control of genome data. We discuss

how statistics computed on iLoci reflect various characteristics of genome content and organization, and illustrate how this can be used to establish a baseline context for evaluating a new genome assembly and annotation. We also introduce a well-defined measure of relative genome compactness, and more generally show how iLoci reveal the extent of gene clustering genome wide.

Conclusions: The iLocus framework, available as open source software as part of the AEGeAn Toolkit (https://brendelgroup.github.io/AEGeAn), provides a comprehensive solution to common challenges in annotation and analysis of non-model genomes.

Background

The ready availability of Next-Generation Sequencing (NGS) technologies has resulted in genome data for thousands of species, with no slowing down of data accumulation in sight. Given this volume of data, fast and accurate computational approaches are needed now more than ever to process the initial sequence data into meaningful units of knowledge about the sequenced genome. The conventional paradigm for that task from a few years ago is outdated. At that time one could expect community groups to carefully assemble and annotate the genomes of their expertise, resulting over a period of time in gap-filled assemblies and refined documentation of genome content in terms of protein-coding genes, non-coding RNA genes, transposable elements, repetitive sequences, and so forth. Such time-consuming and expensive efforts are impractical for the organisms currently being sequenced with NGS technologies.

Out of necessity, the old paradigm has for the most part been replaced by an implicit new standard: genome data are presented as massive short read collections available from databases like the NCBI Sequence Read Archive [102] and in processed form as sets of assembled and computationally annotated scaffolds. Concomitantly, downstream analyses of these data have to be adjusted to scope and quality limitations intrinsic to the new data production process. First, assembly completeness will vary depending on the degree of read coverage and genome complexity (size and repetitiveness). Typically, assemblies will consist of tens to hundreds of large scaffolds, which in the best case can be ordered into linkage groups that approach pseudo-chromosomes; and in addition, manifold more short scaffolds, typically unplaced relative to any linkage groups. Second, annotation will commonly not have been expertly curated, but rather have resulted from first-pass outputs of annotation workflows such as AUGUSTUS [103], MAKER-P [104], BRAKER1 [105], or NCBI Gnomon [106].

Another challenge can be the more temporary nature of the data. As additional sequences can often be acquired cheaply and easily for a species (for example, genomic DNA reads for libraries of different insert sizes; RNA-Seq reads from transcriptome studies under various conditions; or spliced alignments of protein sequences from a newly annotated, closely related species), both the species' genome assembly and its genome annotation may change. However, in the common scenario laid out above, the additional analyses will typically come without the community support to carefully sort out and document all the changes. Thus, over a short span of several years, there may be several annotation versions even for a single stable genome assembly, and it becomes difficult to track references to particular genes and genome features. A pertinent example from our experience is provided by the number of concurrent annotations in recent use for the honey bee (Apis mellifera) genome [107, 108, 109].

How then should one compare results of a study on a current genome assembly and annotation version with previous results in the literature that used a prior assembly/annotation pair? How could one derive subsets of just those gene models that are solidly supported by evidence, to the extent that future genome-wide assembly/annotation improvements will not invalidate these current models? How does one disentangle artifacts of incomplete or inaccurate assembly/annotation from genuine species-specific genome features?

A solution to the problem must address both reproducibility of analyses on genome data and scalability to accommodate thousands of genomes, each potentially with multiple assemblies and annotations. At the core of a solution must be the ability to distinguish what has changed and what has remained invariant from one assembly/annotation pair to another. Discriminating between solid, reliable annotations and annotations of uncertain quality is also crucial in order to enable separation of technical artifacts from effects of interest rooted in the underlying genome biology. Typical examples of this challenge include annotation of UTRs, ncRNA genes, or identification of transposable elements: comparing two genome annotations, one would like to know whether differences in UTR lengths or ncRNA gene and transposon content are due to insufficient data for annotation, annotation workflow settings, or genome evolution.

The ParsEval software [110] provides a convenient tool for comparing two sets of annotations for the same genome assembly. Here we introduce a more general concept and associated software applicable to both single genome analyses and comparisons across assemblies, annotations, and genomes. The basic idea is to represent a given assembly/annotation pair as a set of distinct units that can be largely independently characterized and updated. We show how the parsing of a genome into such distinct iLoci provides a suitable "coordinate system" for working with rapidly changing genome assembly/annotation data. Applications to genome project data for various plant and animal species demonstrate how iLoci analyses can give insights into genome organization and features, as well as assembly and annotation status.

Methods

Toolkit design

Motivated by the challenges of present day genome data reviewed in the **Back-ground** section, we have developed a toolkit for the **A**nalysis and **E**valuation of **Ge**nome

Annotations (AEGeAn [111]). The design of the toolkit followed general principles to achieve reproducible and scalable applications that are easy to use, available as open source code, and integrated with existing tools such as GenomeTools [112, 91]. From a user perspective, the toolkit is meant to work with well-defined inputs consisting of one or more genome sequences paired with associated genome annotation, provided in multi-FASTA and GFF3 [90] formats, respectively. As discussed in the Background section, an input pair can represent a mature model organism genome/annotation version or an incomplete assembly with preliminary annotation. Either way, the toolkit will allow the user to probe genome assembly content and organization, with results reflecting both underlying genome features and the degree of assembly/annotation completeness and accuracy. We show how comparison of different data sets suggests interpretation of results that distinguishes the two possibilities.

The current implementation of the AEGeAn Toolkit provides summary statistics covering a large range of specific questions concerning genome content and organization as well as utility functions to select subsets of genome features for further analysis. Before discussing algorithmic and programming details, we list a number of specific questions that AEGeAn tools address. The **Results** section demonstrates the usefulness of the tool in practical applications.

A first range of questions addresses genome content: How many genes are annotated for a particular assembly/annotation pair? What proportion of the genome is occupied by these genes? What can be said about their length, number of exons, nucleotide composition, and other characteristics? What fraction of genes are protein-coding versus non-coding RNA genes? How many of the gene models have support from transcript evidence, and how many genes can be identified as likely homologs of genes in other species? As we will illustrate later, these seemingly simple questions actually require very precise processing of the annotation file to be reproducibly and meaningfully answered. In particular, the handling of alternative transcription as well as overlapping gene mod-

els needs to be unambiguously defined. The AEGeAn Toolkit includes functions that subselect gene loci based on user-defined characteristics. These functions facilitate the generation of reliable data sets for applications such as codon usage statistics, training of gene prediction models, or identification of transcription regulatory motifs.

A second range of questions addresses genome organization: How densely or sparsely packed are the genes? Is there clustering of genes, and if so, how large are these clusters and what types of genes occur in clusters? More generally, how is the intergenic space organized?

Lastly, all of the above questions are of interest in a comparative genomics context. To what extent are genomes within a clade of species similarly organized? And, maybe even more intriguingly, to what extent is genome organization functionally important?

Conceptual definition of interval loci

To address the toolkit design prescriptions, we introduce a precise parsing of an assembly/annotation pair into smaller units, termed *interval loci*, that provide a robust, granular, and dynamic data set for answering the biological questions posed above. Each interval locus (or *iLocus*) is intended to capture the local genomic context of a genic or intergenic space, providing an alternative coordinate system to the conventional scaffold-based system that is robust to changes in assemblies and annotations. Conceptually, an iLocus is a genomic interval, the boundaries of which are computed from annotated gene models, with an extension to include probable adjacent *cis*-regulatory regions. The precise procedure for computing iLoci is described in detail in the next section.

iLoci can be distinguished by various characteristics, as summarized in **Figure 4.1**. iLoci containing genes are referred to as giLoci, with those encoding protein-coding genes labeled as piLoci and those containing non-coding genes labeled as niLoci. piLoci harboring multiple overlapping gene models are designated complex (ciLoci), while those with a single isolated gene model are designated simple (siLoci). iLoci containing no

gene models are designated as intergenic (iiLoci) if they are flanked on both sides by genes, or as incomplete fragments (fiLoci) if they are are flanked on either side by an end of the scaffold.

To illustrate these concepts, **Figure 4.2** shows the parsing of a hypothetical scaffold into its constituent iLoci. The parsing captures an intuitive and practical decomposition of the genome. The piLoci comprise a superset of protein-coding genes when reporting gene number or calculating descriptive statistics on gene features. However, more reliable results would be expected from the siLoci, or even better a subset of the siLoci with well supported gene models. The ciLoci will typically require a whole lot more attention to establish whether the overlapping gene models reflect observed transcription or are artifacts of unresolved annotation conflicts.

Operational definition of iLoci

Computing iLoci for an assembled contig/scaffold/pseudo-chromosome S depends on a set of intervals G (corresponding to gene models annotated on S) and an extension parameter δ . The basic procedure is described in **Algorithms 1** and **2**. In brief, the ComputeLoci algorithm computes a new set of intervals L such that any two overlapping elements $g_m, g_n \in G$ are bounded by the same interval $loc \in L$. Although the algorithm is general, here g_m and g_n refer to gene bodies, defined as the interval from the start to the end of the respective annotated transcription events. The Extendintervals algorithm then assesses each pair of adjacent intervals $loc_m, loc_n \in L$ and determines how far the intervals can be extended toward each other and whether any additional space remains between them for the creation of a third interval: if the number of nucleotides separating the two intervals $dist(loc_m, loc_n) > 3\delta$ nucleotides, then loc_m and loc_n will be extended toward each other by δ nucleotides, each designated as a giLocus, and the remaining space between them will be designated as an iiLocus; if $2\delta < dist(loc_m, loc_n) \le 3\delta$, then loc_m and loc_n are extended toward each other equally

until they meet, with extensions potentially as long as 1.5δ , to prevent annotating an extremely short iiLocus; if $dist(loc_m, loc_n) \leq 2\delta$, loc_m and loc_n will each be extended by δ resulting in overlapping iLoci. In the two latter cases the toolkit records a zero-length iLoci (ziLocus) between the adjacent giLoci for consistency and calculation of cumulative statistics described below.

Algorithm 1 Compute giLocus boundaries

```
1: procedure Overlap(loc, G)
        O \leftarrow loc
 2:
        for q' \in G do
 3:
             if q' overlaps with loc then
 4:
                 O \leftarrow O \cup g'
 5:
                 mark q' as visited
 6:
 7: return O
 8: procedure ComputeLoci(G, \delta)
 9:
        L \leftarrow \emptyset
10:
        for interval q \in G do
             if q is marked as visited then
11:
                 continue
12:
            interval loc \leftarrow g
13:
             \operatorname{mark} q as visited
14:
             while Overlap(loc, G) \supset loc do
15:
                 loc \leftarrow OVERLAP(loc, G)
16:
             L \leftarrow L \cup \{loc\}
17:
        EXTENDINTERVALS(L, \delta)
18:
19: return L
```

The iLocus parsing procedure is designed with the canonical case of gene organization in mind: a single gene model flanked on both sides by hundreds or thousands of nucleotides of intergenic space. All eukaryotic genomes have exceptions to this case, some to a greater extent than others. The basic parsing procedure can handle some exceptions, such as genes separated by very little intergenic space, but there are additional exceptions that occur frequently enough to merit additional post-processing and refinement.

The basic procedure places two gene models in the same iLocus if their gene bodies

Algorithm 2 Extend giLocus boundaries, identify iiLoci

```
1: procedure ExtendIntervals(L, \delta)
         for adjacent intervals x, y \in L do
 2:
 3:
             if dist(x,y) < 2\delta then
                  End(x) \leftarrow End(x) + \delta
 4:
                  Start(y) \leftarrow Start(y) - \delta
 5:
             else if 2\delta < dist(x,y) < 3\delta then
 6:
                  midpoint \leftarrow |Average(End(x), Start(y))|
 7:
                  End(x) \leftarrow midpoint
 8:
                  Start(y) \leftarrow midpoint + 1
 9:
10:
             else
                  End(x) \leftarrow End(x) + \delta
11:
                  Start(y) \leftarrow Start(y) - \delta
12:
                  interval iiLocus \leftarrow [End(x) + \delta + 1, Start(y) - \delta - 1]
13:
                  L \leftarrow L \cup \{iiLocus\}
14:
```

have any overlap. While this is intended to capture gene models that may be conflicting or misannotated and in need of additional attention to resolve coordinates, an unintended consequence is the occasional grouping of genes with a trivial amount of incidental overlap. For example, if two genes—each a few kilobases in length—happen to have 10-20 nucleotides of overlap in their UTRs, they should be separated and handled as distinct loci. In post-processing, we enable splitting of such trivially overlapping iLoci by introducing two additional parameters: ω , the number of nucleotides that two gene models must overlap to remain in the same iLocus, and κ indicating whether that overlap is calculated using entire gene bodies ($\kappa = 0$) or just the coding sequences ($\kappa = 1$).

The initial procedure also groups non-coding RNA genes and protein-coding genes together if they overlap. In post-processing, ncRNA genes and protein-coding genes are treated separately and will not be grouped in the same iLocus regardless of overlap, although overlapping ncRNA genes are grouped in the same niLocus.

An additional exception occurs when a gene resides completely within a single intron of another gene. These genes are placed in the same iLocus during the initial parsing procedure, but can be separated into distinct iLoci during post-processing.

Implementation

In keeping with the conventions implemented by the GenomeTools library [112], most of the core functionality of the AEGeAn Toolkit [111] is implemented by means of node streams for sequential processing of genome features that are represented as feature graphs. In brief, genome features such as genes, exons, UTRs, and coding sequences are represented as nodes in a directed acyclic graph, and parent/child relationships between features, denoted by ID and Parent attributes in GFF3, are represented as edges in the graph. Each connected component (CC) in the graph, typically corresponding to a gene and its subfeatures, is then processed sequentially by one or more node streams, each designed for a specific annotation processing task. One advantage of this approach is its low memory footprint, as at most only a small number of CCs need be loaded into memory at any given moment.

The AgnLocusStream module in the AEGeAn Toolkit implements a node stream for computing iLocus boundaries. This node stream expects as input gene annotations (CCs with a gene feature as the root node) sorted by genomic position, but it is designed to work with arbitrary feature types. Initially, the node stream will collect a single gene feature from the input and store it in a buffer. Then, as subsequent gene features are collected, they are tested for overlap with the current iLocus (buffer) and accumulated as long as their leftmost position is less than or equal to the rightmost position of the buffer. When the node stream encounters a gene feature that does not overlap with the buffer, a giLocus feature is created, all the genes in the buffer are assigned as children to the giLocus, a reference to the giLocus is stored temporarily in the node stream, and the giLocus is emitted for further processing or storage. Later, as the subsequent giLocus is emitted, the adjacent pair of giLoci are extended toward each other and, when appropriate, an iiLocus is created and emitted, as described in Algorithm 2.

The AEGeAn Toolkit's AgnLocusRefineStream module implements a node stream for post-processing the initial iLocus designations, as described in the previous section. Any

genes belonging to the same giLocus that do not overlap by at least ω nucleotides in their gene bodies (or coding sequences if $\kappa = 1$), as well as genes contained completely within the intron of another gene, are split into distinct overlapping giLoci.

The *LocusPocus* program is a command-line tool included in the AEGeAn Toolkit distribution which provides a user interface to the *AgnLocusStream* and *AgnLocusRe-fineStream* modules. A detailed description of command-line usage and program inputs and outputs is provided in the **Supplementary Methods**.

Genome content statistics

As discussed in the **Background** section, describing genome characteristics and comparing across species requires selection of reliable subsets of data for analysis. The precise selection criteria used will depend on the questions being asked, but commonly involve a small set of descriptive statistics that can easily be computed from the iLocus sequence and/or associated annotation. These include the length and nucleotide composition of the iLocus itself, as well as the count, length, and composition of corresponding features such as genes, RNAs, exons, introns, and coding sequences. Statistics are computed by invoking the *stats* task of the *genhub-build.py* script (see **Supplementary Methods**) and are stored in tab-separated plain text (.tsv) files to facilitate importing into popular statistical computing packages.

Additional characteristics, such as the quality and homology status of gene models, are not always readily accessible from the iLocus sequence or annotation but can be computed and attached to an iLocus annotation using key-value pairs in GFF3's attribute column. Annotation quality can be measured with statistics such as Maker's annotation edit distance [6] or the GAEVAL integrity score [113], and homology status can be determined via BLAST search or clustering of iLocus protein products. Attaching these characteristics to the iLocus provides additional features on which to filter data.

Descriptive statistics are reported only for a single annotated transcript at each iLo-

cus to ensure aggregate statistics are not biased by redundancy in the data from, for example, genes with many annotated isoforms. The reported transcript is selected according to the amino acid length of its translation product: the transcript with the longest product is reported. In cases where multiple transcripts have translation products of identical length, the transcript with the lexicographically smallest **ID** attribute is reported, ensuring reproducible and deterministic reporting.

Cumulative lengths of different iLocus types are calculated after proper accounting of any iLocus overlaps to ensure each nucleotide in the genome is counted only once (see **Supplementary Methods**). When reported as a fraction of the entire genome, the genomic space occupied for different iLocus categories is calculated as a fraction of effective genome size—the total number of nucleotides in the genome that do not reside within fiLoci—to mitigate potentially confounding inflation of genome size by many short unannotated sequences or sequence fragments.

Genome organization statistics

Beyond genome content, the iLocus framework also allows to systematically study different aspects of genome organization. Here we focus on gene orientation and spacing: are there species-specific patterns of gene arrangements, and how do natural genomes differ in these respects from statistical expectation? Because of the flexible design of the code base described in the **Implementation** section, these questions can easily be generalized and extended, for example with respect to selection of subtypes of genic loci.

To study gene orientation, the *LocusPocus* program reports for each iiLocus the orientation of the flanking giLoci as FF (forward >>), RR (reverse <<), RF (outward <>), or FR (inward ><). In the case that an iiLocus is flanked by one or more ciLoci, the orientation of the gene models directly flanking the intergenic space are reported. These data are used to identify the longest stretches of genes all on the forward strand, all on the reverse strand, or periodically alternating between strands. Differences in

occurrence numbers and lengths of outward and inward iiLoci are also determined for possible interpretation in terms of promoter architecture.

Long iiLoci are flagged as regions for annotation review. More generally, for each giLocus the lengths of the flanking iiLoci are reported. In cases where a giLocus abuts or overlaps with another giLocus, the corresponding iiLocus length is set to zero, and the number of overlapping nucleotides is recorded. The software tracks these cases as zero-length iiLoci (ziLoci). The iiLoci lengths are used in two different ways to reveal gene spacing characteristics. The distribution of aggregate lengths of n adjacent iiLoci shows the mode of typical gene spacings as well as outliers. Secondly, overlapping or abutting giLoci are collapsed into merged iLoci (miLoci) during post-processing and represent gene clusters. Our results show that relative counts and lengths of miLoci compared to non-merged siLoci provide robust measures of genome compactness.

To evaluate observed gene spacing patterns with statistical expectation, we implemented a procedure to generate randomized gene arrangements relative to the input genome annotation in GFF3 format. First, iLoci are computed with $\delta=0$ to identify the precise boundaries of annotated genic regions. Next, giLoci are removed from the sequence and the remaining iiLoci are concatenated. Then, new positions are randomly selected from a uniform distribution for re-inserting the giLoci in shuffled order into the sequence. As each giLocus is re-inserted, the genomic sequence is expanded, and all downstream re-insertion sites are adjusted accordingly. Re-running the iLocus parsing procedure and computing neighbor statistics on these random arrangements provides a baseline for comparison, revealing how genome annotations as observed differ (at the genome scale) from what could be expected from a completely random arrangement of genes.

Comparing assembly/annotation pairs: iLocus stability

Given two assembly/annotation versions for the same genome, we evaluate the stability of iLoci across the two versions using sequence alignment and interval overlap. Specifically, iLocus sequences from the earlier assembly/annotation version (version A) are aligned to the genome of the newer version (version B), and the interval of each alignment is compared to the intervals of iLoci annotated on B. An iLocus from A is designated stable if the following criteria are satisfied: it aligns to genome B with at least 95% identity; an iLocus from B overlaps with at least 90% of the aligned iLocus sequence from A; and the alignment interval overlaps with at least 90% of that same iLocus from B.

For this study, we used Vmatch [114] to compute iLocus alignments (using options—complete—e 5b—identity 95—d—p—showdesc 0). As a preliminary step, we used RepeatMasker [115, 116] and bedtools [117] to identify and filter out iLoci with substantial repetitive content to make alignment computations tractable. Specifically, any iLocus with at least 500 bp or 25% repetitive content was excluded from the Vmatch alignment procedure.

Comparing genome content and organization between related genomes: homologous iLoci

Given a set of annotated genome assemblies for a clade of related species, we compute homologous iLoci (hiLoci) via a protein clustering procedure. For each species, a representative protein sequence is selected for each piLocus (as described in the **Genome content statistics** section). The distinct protein complements from all species are then combined, and the aggregate collection of protein sequences is clustered using cd-hit [118].

In brief, cd-hit processes proteins iteratively from longest to shortest. The first protein is assigned to a cluster by itself and is designated the *representative sequence* of the

cluster. Each subsequent protein is compared to all previous clusters: if the alignment of the protein to a cluster's representative sequence satisfies the specified sequence identity, length similarity, and alignment coverage criteria, it is added to that cluster, and the program advances to the next protein; if a protein cannot be added to any cluster by user-specified clustering criteria, it is placed in a new cluster by itself and designated the representative sequence of that cluster.

Following the clustering procedure, a homologous iLocus (hiLocus) is designated for each protein cluster and the piLoci corresponding to the proteins in that cluster are assigned to that hiLocus. The hiLocus thus provides a link between piLoci from related species and a relative measure of how well conserved the corresponding protein is within the given clade.

This protein clustering procedure is invoked using the *cluster* task of the *genhub-build.py* script. The script provides the following default clustering criteria, although these defaults can be overridden and additional criteria set by the user. The default parameters are as follows: sequence identity $\geq 50\%$; length difference $\leq 65\%$; alignment coverage for longer sequence $\geq 75\%$; alignment coverage for shorter sequence $\geq 85\%$. On the command line these parameters are specified as -c = 0.50 - s = 0.65 - aL = 0.75 - aS = 0.85.

Data sets analyzed

To demonstrate the utility of iLoci for providing a descriptive overview of genome composition and organization, we retrieved RefSeq genome assemblies and corresponding annotations for ten model organisms (complete species names listed shown **Table 4.1**). Species were selected to provide a broad sampling of eukaryotic diversity, with a preference for robust model organisms with mature chromosome-level genome assemblies and extensive community-supported annotation. For each species, we computed iLoci and associated feature statistics, including length, nucleotide composition, exon count,

and effective length using standard GenHub build tasks as described before. Precise configurations and commands run are described in detail in the **genome-summary** section at https://github.com/BrendelGroup/IntervalLoci.

Using iLocus summaries of these ten model organisms as a baseline for comparison, we characterized the genome content and organization of three additional species of interest that serve as important experimental models for evolutionary and ecological studies. The alga *Volvox carteri* is the focus of research on the evolution of cellular complexity and development, and was the second alga (after *Chlamydomonas reinhardtii*) to have its genome sequenced. The paper wasp *Polistes dominula* is an important model for studying the evolution of social behavior, and provided one of the first sequenced genomes of a vespid wasp [119]. The water flea *Daphnia pulex* is a species of ecological and evolutionary interest, and was the first crustacean to have its genome sequenced [120]. Genomes for these three species were processed using the same procedure as the ten model organisms (see https://github.com/BrendelGroup/IntervalLoci).

To investigate the extent of gene conservation in the green algae (phylum: Chlorophyta), we collected and processed data for nine chlorophyte species (Auxenochlorella protothecoides, Chlamydomonas reinhardtii, Chlorella variabilis, Coccomyxa subellipsoidea, Micromonas pusilla, Micromonas sp. RCC299, Ostreococcus lucimarinus, Ostreococcus tauri, and Volvox carteri), as well as four land plants (Arabidopsis thaliana, Brachypodium distachyon, Medicago truncatula, and Oryza sativa) as an outgroup. Retrieval of annotations and sequences and calculation of hiLoci was invoked using standard procedures as described before (see chlorophyta section at https://github.com/BrendelGroup/IntervalLoci). Following the protein clustering procedure, each hiLocus was assigned a preliminary classification: highly conserved if it had a representative from at least four chlorophyte genomes; conserved if it had a representative from at least four chlorophyte genomes; matched if it had a representative from at least two genomes (including the outgroups); and unmatched if it had a representative from only a single genome.

Table 4.1 iLocus content of genomes from ten model organisms and three additional species.

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Species	$\mathrm{Mb^1}$	$\#\mathrm{Seq}^2$	fiLoci	iiLoci	niLoci	siLoci	ciLoci
Saccharomyces cerevisiae	12.1	16	11	289	393	5,704	90
$Cae nor hab dit is\ elegans$	100.3	6	4	7,783	19,230	20,071	81
$Chlamydomonas\ reinhardtii$	120.2	1,556	1,487	6,248	0	14,254	42
$Medicago\ truncatula$	412.8	2,186	2,325	36,465	1,075	50,444	0
$An opheles\ gambiae$	265.0	8,089	8,041	7,724	639	12,184	120
$Drosophila\ melanogaster$	143.7	1,869	1,849	3,436	3,289	13,115	239
$Xenopus\ tropicalis$	1437.5	7,727	8,004	18,580	5,199	21,704	135
Danio rerio	1371.7	1,060	1,276	23,978	12,776	26,229	166
$Mus\ musculus$	2725.5	21	42	23,689	14,103	21,305	218
$Homo\ sapiens$	3088.3	24	48	22,242	16,995	19,067	399
Volvox carteri	137.7	1,251	1,198	7,790	0	14,346	44
$Polistes\ dominula$	208.0	1,483	1,697	5,970	338	11,376	204
$Daphnia\ pulex$	197.3	5,191	4,759	13,052	0	30,456	158

¹Total number of nucleotides in the genome assembly.

iLoci initially classified as *unmatched* were subjected to additional screening to distinguish conserved proteins lacking a nearly-full-length match (due to incomplete or incorrect annotation, or true evolutionary divergence) from orphan proteins without any reliable match. iLoci with a BLASTP match against another species (-evalue 1e-10) were reclassified as *matched*, while those lacking a match were reclassified as *orphan*.

Results

iLoci provide an informative decomposition of genome content

We computed iLoci for ten model organisms covering a wide range of eukaryotic diversity, and provide a summary of each genome and its iLocus complement in **Table 4.1**. The genome assembly sizes in this sampling of eukaryotes spans three orders of magnitude, ranging from 12.1 Mb in *Saccharomyces cerevisiae* to over 3 Gb in *Homo sapiens*. Several genomes are represented exclusively by chromosome sequences, some exclusively by unplaced genomic scaffolds, and some by a combination of both. The

²Total number of assembled (pseudo-)chromosomes or unplaced genomic scaffolds or, for some species, both.

Table 4.2 Summary of piLoci from genomes of ten model organisms and three additional species.

species.			
Species	piLoci	$Occupancy^1$	Single Exon piLoci
Saccharomyces cerevisiae	5,794	11.4 Mb (94.3%)	5,529 (95.4%)
$Cae nor hab ditis\ elegans$	20,152	71.2 Mb (71.0%)	579~(~2.9%)
$Chlamydomonas\ reinhardtii$	14,296	74.1 Mb (68.2%)	1,127 (7.9%)
$Medicago\ truncatula$	50,444	179.5 Mb (44.9%)	$11,481\ (22.8\%)$
$An opheles\ gambiae$	12,304	83.7 Mb (35.9%)	$1{,}137~(~9.2\%)$
$Drosophila\ melanogaster$	13,354	95.1 Mb (70.0%)	$2,062 \ (15.4\%)$
$Xenopus\ tropicalis$	21,839	686.6 Mb (50.4%)	1,353~(~6.2%)
Danio rerio	26,395	793.7 Mb (58.6%)	1,062~(~4.1%)
$Mus\ musculus$	21,523	1,014.1 Mb (38.2%)	$2,326 \ (10.8\%)$
$Homo\ sapiens$	19,466	1,204.7 Mb (40.0%)	$1,\!270~(~6.5\%)$
Volvox carteri	14,390	89.2 Mb (69.1%)	1,086 (7.5%)
$Polistes\ dominula$	11,580	64.1 Mb (34.8%)	455 (3.9%)
$Daphnia\ pulex$	30,614	89.2 Mb (54.3%)	5,053 (16.5%)

¹Total number of nucleotides occupied by piLoci, and the corresponding fraction of effective genome size.

number of fiLoci, with a strict upper bound of twice the number of assembled sequences, is informative primarily with respect to assembly status. For most of these genomes the actual number of observed fiLoci is close to half of the upper limit, the primary contributing factors being the presence gene annotations near the ends of the genomic sequences and the presence of many unannotated scaffolds.

iiLoci correspond to intergenic DNA and are reflective of genome organization. There can be at most n-m iiLoci in a genome with n genes and m annotated sequences, but closely-spaced genes will reduce the number of observed iiLoci, as will the presence of unannotated scaffolds.

Differences in the number of niLoci appear to reflect the varying extent to which noncoding RNA genes annotated by NCBI and the respective model organism communities have been integrated into RefSeq, rather than the actual relative abundance of these genes in the respective genomes.

The abundance of piLoci in each genome (representing distinct protein-coding regions) spans just a single order of magnitude, from 5,794 piLoci in *Saccharomyces cerevisiae* to 50,444 in *Medicago truncatula* (**Table 4.2**). The total space occupied by piLoci,

Table 4.3 Summary of miLoci from genomes of ten model organisms and three additional species.

tional species.				
Species	miLoci	$Occupancy^1$	Gene Count ²	$Singletons^3$
Saccharomyces cerevisiae	240	11.4 Mb (95.0%)	15	65 (1.1%)
$Cae nor hab dit is\ elegans$	5,178	70.4 Mb (70.2%)	4	2,611~(~6.6%)
$Chlamydomonas\ reinhardtii$	3,042	54.9 Mb (50.5%)	3	$3,753 \ (26.3\%)$
$Medicago\ truncatula$	9,799	81.2 Mb (20.3%)	2	27,702 (53.8%)
$An opheles\ gambiae$	2,198	33.9 Mb (14.5%)	2	5,962 (46.1%)
$Drosophila\ melanogaster$	1,867	88.3 Mb (65.0%)	4	1,621 (9.7%)
$Xenopus\ tropicalis$	3,489	206.8 Mb (15.2%)	2	$17,009 \ (62.9\%)$
Danio rerio	5,752	285.4 Mb (21.1%)	2	18,834 (48.1%)
$Mus\ musculus$	6,625	746.9 Mb (28.1%)	2	17,085 (48.0%)
$Homo\ sapiens$	$7,\!222$	1,033.9 Mb (34.3%)	2	15,044 (41.3%)
Volvox carteri	3,240	57.5 Mb (44.6%)	2	5,210 (36.2%)
$Polistes\ dominula$	2,323	36.1 Mb (19.6%)	3	$3,960 \ (33.2\%)$
$Daphnia\ pulex$	$6,\!284$	59.2 Mb (36.0%)	3	$9,765 \ (31.9\%)$

¹Total number of nucleotides occupied by miLoci, and the corresponding fraction of effective genome size.

however, spans three orders of magnitude, similar to genome size. This is explained by a distinct contrast in siLocus length between vertebrates and the other species (**Figure S1**), the compound result of increases in both intron abundance and length (**Figures S2-S3**). We note that while the protein-coding portion of the human genome is commonly reported as 2-4%, this refers only to protein-coding exons. The inclusion of introns and UTRs places the protein-coding fraction of the genome at approximately 40% for both human and mouse.

ciLoci occur in the dozens or hundreds in most genomes, accounting for only a small proportion of genes. Only one genome from these ten species, that of *Medicago truncatula*, harbors no ciLoci.

iLoci reflect patterns of genome organization

Gene clustering is abundant in eukaryotic genomes

Although there are examples of well-described gene clusters, such as those associated with *Hox* genes [121], the spatial distribution of genes in general is not well characterized

²Median per-miLocus gene count per miLocus. Histograms showing full distributions available in **Figure S4**.

³Total number of giLoci not contained in a miLocus, and corresponding fraction of all giLoci.

in eukaryotes. Hox clusters are composed of functionally related developmental genes with a conserved colinear arrangement, a common direction of transcription, and close proximity in the genome. More generally, gene clusters described in the literature need not be comprised of only genes that are directly adjacent, but are loosely defined as sets of genes of a common function situated much closer to each other than would be expected by chance [122].

However, the extent to which genes are tightly packed throughout the entire genome and the characteristics of these gene-dense regions are not well defined or understood in eukaryotes. *Merged iLoci (miLoci)* provide a well-defined unit of analysis for investigating the spatial distribution of genes genome-wide. Using miLoci, we surveyed genome organization in ten model organisms.

Genes cluster together frequently in eukaryotic genomes. The most frequent groupings involve a small number (2-4) of genes (see **Table 4.3** and **Figure S4**), but all genomes include larger clusters involving dozens or even hundreds of tightly packed genes. The budding yeast *Saccharomyces cerevisiae* is an extreme example, populated almost entirely by just 240 miLoci encompassing all but 65 genes in the entire genome. *Caenorhabditis elegans* and *Drosophila melanogaster* also bear signatures of a higher overall level of genome compactness, with larger numbers (and overall proportion) of genes merged into miLoci and a larger proportion of genomic sequence occupied by miLoci.

In general, clustered genes do not differ substantially in length or nucleotide composition from spaced out genes (**Figure S5**). However, especially among large miLoci, clustered genes are often functionally related. The longest miLoci in the human genome include a cluster of 23 snoRNA genes, a cluster of V(D)J adaptive immunity genes, and a cluster of keratin associated proteins, while in mouse the longest miLocus is comprised of 72 microRNA genes. In the non-mammal vertebrates, the longest miLoci are comprised exclusively of long stretches of hundreds tRNA gene annotations. As tRNA-derived SINE

transposons are known to be abundant in at least one of these species [123], and no annotations for such transposons appear to be included in the RefSeq annotation, it is likely these miLoci capture large clusters of misannotated repetitive elements. Within the two plant genomes studied (*Medicago* and *Chlamydomonas*) miLoci include a large proportion of genes of unknown function (encoding "hypothetical" and "predicted" proteins), although ribosomal proteins, membrane proteins, photosystem II proteins (in *Medicago*), and flagellar proteins (in *Chlamydomonas*) feature prominently in the largest miLoci.

The spacing of genes over longer ranges is revealed by distributions of aggregate lengths of n adjacent iiLoci. Long-range spacing of genes varies considerably in eukary-otes, with some species exhibiting homogenous gene spacing over relatively short spans (spans of 5-10 genes in *Caenorhabditis elegans* and *Medicago truncatula*), and others showing heterogenous spacing even over long spans (spans of more than 30 genes in *Mus musculus*; see **Figure 4.3**).

Compactness of eukaryotic genomes varies widely

Figure 4.4 further explores the notion of compactness of a genome by two complementary measures calculated on the constituent chromosome or long scaffold sequences: σ , defined as the proportion of the sequence occupied by miLoci; and ϕ , defined as the fraction of giLoci in the sequence merged into miLoci. Distinct quadrants in the plot reflect characteristic overall genome organization. Low values of ϕ associated with low values of σ (lower left) correspond to genes as "islands" in an "ocean" of intergenic (presumably repetitive) DNA. High values of ϕ associated with low values of σ (lower right) correspond to "archipelagos" of genes. And high values of ϕ associated with high values of σ (upper right) correspond to "compact" genome organization.

Let the average iiLocus length be ρ -times the average giLocus length. Then it is easy to show that

$$\sigma \approx \frac{\phi}{1 + (1 - \phi)\rho}.$$

Figure 4.4 gives the curves for ρ equal to 0.1, 1, 2, and 4.

Empirical (ϕ, σ) values calculated for continuous genome sequences of at least 1 Mb for the 10 model species reveal a wide range of genome compactness across eukaryotes, yet show remarkable consistency within species (**Figure 4.5**) and even within clades (as confirmed by sampling of additional species within 4 clades; see **Figure S5**). Genome compactness scales roughly with genome size, at least across major clade divisions and levels of organismal complexity. Within Chlorophyta, compactness scales almost perfectly with genome size, although this trend is not maintained in clades characterized by larger genome sizes. Sequences from *Saccharomyces cerevisiae* are the most compact of all 10 model organisms analyzed, consistent with the paucity of iiLoci in the genome and the highest gene counts per miLocus observed in these data sets.

Very few sequences show extremely low levels of compactness: only six sequences have $\phi < 0.2$ and $\sigma < 0.2$, two of which correspond to mammalian sex chromosomes, with the other four corresponding to unplaced scaffolds from *Xenopus tropicalis*. Likewise, very few sequences are dominated by an "archipelago"-type organization (high ϕ and low σ). Those with $\phi > 0.7$ and $\sigma < 0.3$ are annotated almost exclusively with long stretches of dozens or hundreds of tRNA gene annotations in *Xenopus tropicalis* and *Danio rerio*.

Gene clustering occurs more frequently than expected by chance

The framework provided by miLoci for investigating genome organization enables us to investigate whether the amount of clustering observed in a particular genome assembly/annotation pair is more or less than would be expected by chance. For long chromosome and scaffold sequences (\geq 1Mb) in ten model organisms, we computed random spatial arrangements of the genes on each sequence and then re-computed iLoci and associated summary statistics for comparison with the observed annotation.

Random positioning of genes results in decreased amounts of gene clustering in most species, as reflected by several features: a decrease in the number of miLoci; a decrease

Table 4.4 Summary of miLoci computed on randomly positioned genes.

	v	1	√ 1	0
Species	miLoci	Occupancy	Gene Count	Singletons
Saccharomyces cerevisiae	134	4.6 Mb (96.7%)	14	10 (0.4%)
$Caenorhabditis\ elegans$	$6,\!389$	71.7 Mb (71.5%)	4	3,186 (8.1%)
$Chlamydomonas\ reinhardtii$	2,502	34.0 Mb (42.7%)	2	$3,866 \ (36.6\%)$
$Medicago\ truncatula$	$9,\!189$	73.8 Mb (19.2%)	2	$27,408 \ (56.0\%)$
$An opheles\ gambiae$	1,506	26.2 Mb (11.4%)	2	$8,982 \ (72.4\%)$
$Drosophila\ melanogaster$	3,873	81.8 Mb (59.5%)	3	$3,960 \ (23.9\%)$
$Xenopus\ tropicalis$	$2,\!193$	161.0 Mb (12.5%)	2	$17,372 \ (77.7\%)$
Danio rerio	4,683	251.8 Mb (18.8%)	2	25,659~(68.4%)
$Mus\ musculus$	5,483	633.1 Mb (23.2%)	2	21,904~(61.5%)
$Homo\ sapiens$	6,210	898.2 Mb (29.1%)	2	$19,834 \ (54.4\%)$

Compare to Table 4.3.

in the space occupied by miLoci; a decrease in the number of genes per miLocus; and an increase in the number of singleton genes not associated with miLoci (**Table 4.4**). These measures are completely consistent for all but the most compact genomes: For example, in *Drosophila melanogaster* we observe an increase in the number of miLoci and yet a decrease in the number of genes per miLocus and the proportion of the genome occupied by miLoci. Every genome has evidence of reduced gene clustering from at least one feature, and most genomes have consistent evidence from every feature.

Random arrangement of genes also influences signatures of genome compactness, reflecting less compactness relative to the actual annotated positioning of genes. The (ϕ, σ) statistics calculated on long genomic sequences are consistently lower for random arrangements than actual arrangements for all ten model species (**Figure 4.6**), with the exception of *Saccharomyces cerevisiae*.

Characterization of three species of interest

With baseline expectations about genome content and organization established by iLocus analysis of ten model organism genomes, we sought to characterize the composition and organization of genomes from 3 additional species: the multicelullar green alga *Volvox carteri*, the primitively eusocial paper wasp *Polistes dominula*, and the microcrustacean *Daphnia pulex*.

The genome content of $Volvox\ carteri$ is very similar to that of $Chlamydomonas\ reinhardtii\ (Tables 4.1-4.3)$. Characteristics of protein-coding regions (summarized in Table 4.2) show particular similarity, as reflected by the number of piLoci, the total space occupied by these piLoci, and the fraction of piLoci corresponding to single-exon genes. Volvox also exhibits a similar, though slightly lower, level of genome compactness to Chlamydomonas: in Volvox, miLoci account for a smaller percentage of the genome sequence and a smaller fraction of the overall gene count (Table 4.3). The (ϕ, σ) values measuring genome compactness fall within a similar range for Volvox and Chlamydomonas, with Volvox shifted to slightly lower values for both measures (Figure S5A). In total, these observations are consistent with the claims that, despite an estimated 50-200 million years of divergence and major differences in cellular complexity, the genomes and proteomes of Volvox and Chlamydomonas are impressively similar [124].

The genome composition of *Polistes dominula* bears some similarities as well as some differences with the model organisms studied. The number of annotated protein-coding regions in *Polistes dominula* and the proportion of genome sequence occupied by these regions is at the lower limit observed in the insects and most other species. Of particular note is the small number single-exon genes in *Polistes dominula*, both in terms of absolute number and percentage of all genes (**Table 4.2**), likely a reflection of strict filtering of single-exon transcript alignments in the annotation of the genome [119]. The organization of the *Polistes* genome tends toward the "archipelago"-style arrangement, with a large proportion of genes merged into clusters that are spaced out and therefore occupy a smaller fraction of the genome sequence (**Figure S5C**). We also observe more variation in genome organization in *Polistes* (and the insects in general) than is observed in the ten model organisms overall. This observation could represent heterogenous gene organization throughout these genomes, but could also be the result of a more fragmented genome assembly (true of many insect genomes) or missing genes in the genome annotation (possible with *Polistes*).

In many ways the profile of the Daphnia pulex genome is intermediate relative to the survey of ten model organisms. The most striking feature of Daphnia is its large number of annotated genes (with a piLocus count that is second only to Medicago) and large fraction of single-exon piLoci (Table 4.2). The proportion of the genome occupied by these genes, however, is average with respect to other arthropods and to the ten model species in overall. Claims regarding the compactness of the *Daphnia* genome, based primarily on average gene spans and average intron lengths, are not supported by our analysis [120]. We confirm that genes are on average shorter in Daphnia than Drosophila, despite a larger number of exons per gene (Figure S1). However, this appears to be influenced more by reduced exon length rather than by intron length: median intron length is almost indistinguishable between Daphnia and Drosophila (75 bp and 72 bp, respectively; see **Figure S3**), whereas exon length is substantially shorter in Daphnia (154 bp and 286 median exon length, respectively; see **Figure S6**). Further, we observe consistently lower (ϕ, σ) values for Daphnia than for Drosophila, reflective of a smaller fraction of tightly-packed genes and a smaller proportion of the genome sequence occupied by such gene clusters (Figure S5C). Across all of these measures, Daphnia pulex is characterized by a moderate level of genomic compactness relative to other arthropods and eukaryotes in general.

iLoci provide a robust representation of the genome

Improvements in genome assemblies come at the expense of disrupting the sequence-based coordinate system typically used for annotating the location of genome features. Parsing an annotated genome into iLoci provides an alternative representation of the genome that is robust to assembly and annotation updates. To assess the stability of iLoci in genome projects over time, we selected two model organisms (Arabidopsis thaliana and the honey bee Apis mellifera) for which multiple assemblies and annotations produced over the span of several years are available. The 2005 TAIR6 release was the

first annotation of the A. thaliana genome managed by The Arabidopsis Information Resource [125], while the 2010 TAIR10 release integrates TAIR's latest improvements to both the reference genome assembly and annotation [126]. Between TAIR6 and TAIR10, changes to the genome sequence were quite limited and were accompanied by a relatively small increase in the number of annotated gene models (see **Table 4.5**).

Table 4.5 Descriptive summary of annotated genome assemblies for A. thaliana and A. mellifera.

тентена.	TAID 6	TAID 10	0.00 4.01	0.00 2.22	
	TAIR6	TAIR10	$OGSv1.0^{1}$	$OGSv3.2^2$	
Assembly size (Mb)	119.2	119.1	228.6	234.1	
Sequences	5	5	7,655	5,644	
Protein-coding genes	26,541	27,202	10,157	15,314	
ncRNA genes	769	1,290	-	-	
Total iLoci	37,299	37,669	22,784	26,251	
siLoci	26,507	27,003	9,898	$14,\!531$	
ciLoci	17	96	106	292	
niLoci	769	1,286	-	-	
iiLoci	9,997	$9,\!276$	4,790	6,160	
fiLoci	9	8	7,990	$5,\!268$	

 $^{^{1}}$ Corresponds to assembly Amel_2.0.

For A. mellifera, the Honey Bee Genome Sequencing Consortium's assembly version Amel_2.0 and Official Gene Set 1 (OGSv1.0) were preliminary data resources in use prior to the initial published description of the honey bee genome in 2006 [107], while assembly Amel_4.5 and OGSv3.2 represent the consortium's latest improvements to the genome and corresponding annotation as of 2014 [108]. In contrast to Arabidopsis, upgrades to the genome assembly between Amel_2.0 and Amel_4.5 were quite substantial, including enriched sequencing of low-%GC-content DNA, as well as additional coverage from 454 and SOLiD reads [108]. Furthermore, between OGSv1.0 and OGSv3.2 there was a dramatic increase in the number of annotated gene models, up from approximately 10,000 to over 15,000 genes (see Table 4.5), with a corresponding drop in the number of exons per gene.

²Corresponds to assembly Amel_4.5.

For both species, we computed iLoci for each assembly/annotation version and determined *iLocus stability* as described in the **Methods**. **Figure 4.7** provides a breakdown of stability by iLocus type. Essentially all iLoci from TAIR6 (33,717 / 33,730) align to the TAIR10 genome, with a large majority (28,890 / 33,730 = 86%) satisfying the criteria to be designated as *stable*. In *Apis mellifera*, we also observe high similarity at the sequence level, with 93% (18,101 / 19,447) of iLoci from Amel_2.0/OGSv1.0 aligning to the Amel_4.5/OGSv3.2 genome. However, only 28% (5,519 / 19,447) of iLoci from OGSv1.0 satisfy the criteria to be designated *stable*. In both species, a higher proportion of *siLoci* are stable than of iiLoci, likely as a result of intergenic spaces being punctuated by additional gene models in the newer annotations.

Figure 4.8 shows the length and %GC Content of siLoci and iiLoci for both species, highlighting characteristic differences between stable and unstable iLoci. There is very little to differentiate stable and unstable siLoci in *Arabidopsis thaliana*, which is unsurprising given the incremental nature of annotation updates between TAIR6 and TAIR10. For *Apis mellifera*, however, we observe much less stability for long siLoci, with only a small percentage of siLoci longer than 10kb designated as stable (Table S2). For both species we observe less stability in long iiLoci, in both cases the likely result of an increase in gene models in previously unannotated regions.

We see very little change in iLocus classification among stable iLoci (see **Table 4.6**). For example, the majority of siLoci remain siLoci: that is, at stable loci a protein-coding gene annotation is rarely replaced with a non-coding gene annotation or no annotation at all. The only minor exception to note is the conversion of 285 fiLoci in OGSv1.0 to siLoci in OGSv3.2 with the addition of new gene models to previously unannotated ends of assembly scaffolds.

Table 4.6 A breakdown of stable iLoci by type. Arabidopsis thaliana (TAIR6 \rightarrow TAIR10)

	-		`		/
	siLocus	ciLocus	niLocus	iiLocus	fiLocus
siLocus	22,717	24	5	3	0
ciLocus	1	11	0	0	0
niLocus	0	0	730	0	0
iiLocus	21	1	0	5,366	0
fiLocus	0	0	0	0	4

Apis mellifera (OGSv1.0 \rightarrow OGSv3.2)

	siLocus	ciLocus	niLocus	iiLocus	fiLocus
siLocus	2,659	15	0	0	23
ciLocus	5	3	0	0	0
niLocus	0	0	0	0	0
iiLocus	13	0	0	554	0
fiLocus	285	0	0	28	1934

Rows correspond to the type annotated in the earlier annotation version, and columns correspond to the type annotated in the later version. For example, 5 stable iLoci annotated as siLoci in TAIR6 were subsequently designated as niLoci in TAIR10.

iLoci reveal extent of gene conservation in green algae

The green algae (phylum Chlorophyta) diverged from land plants an estimated 1 billion years ago [127] and encompass a diverse set of organisms ubiquitous in marine and soil environments. Chlorophytes exhibit substantial variation in physical stature, genome size, and cellular complexity, and include many important systems for study of the evolution of multicellularity and photosynthesis. The publication of the *Volvox carteri* genome [124] reported over 5,000 protein families conserved between *Volvox* (a multicellular alga) and *Chlamydomonos reinhardtii* (a unicellular relative), accounting for over a third of the species' respective proteomes. With nine representative chlorophyte genomes now available from RefSeq [128], we sought to leverage iLoci to characterize the extent of gene conservation throughout the entire phylum and characteristics of conserved and species-specific genes.

Figure 4.9 shows a breakdown of each genome according to iLocus type and conservation status, showing both the number of iLoci in each category as well as the proportion

of the genome occupied by iLoci from each category. Counts and aggregate space occupied by intergenic regions and assembly fragments (iiLoci and fiLoci, respectively) reflect the diversity of genome size and gene density across Chlorophyta. A small number of piLoci from each genome are designated as *orphans*, indicating no reliable protein match in any other species, with the majority designated as *matched*, having at least one match in another species. The designations *conserved* and *highly conserved* are reserved for those piLoci whose protein products are not only well conserved throughout the phylum (*conserved*: conserved in at least 4 species; *highly conserved*: conserved in all 9 species), but also have near-full-length alignments with conserved orthologs.

Given these stricter criteria, we observe approximately 100 highly conserved piLoci and approximately 1,000 conserved piLoci in each species, accounting for only a small proportion of the gene space. Highly conserved piLoci are associated with a variety of cellular components and processes including ribosomal proteins, histone proteins, and kinase/phosphatase activity. Unsurprisingly, the vast majority of orphan piLoci are annotated as "predicted" or "hypothetical proteins". Amongst the handful with functional annotations, flagellar-associated proteins are prominent in Chlamydomonas reinhardtii orphans, while transposition proteins are prominent in Volvox carteri orphans. Conserved and highly conserved piLoci tend to be longer on average and restricted to a smaller range of length and nucleotide composition than orphans (see Figure 4.10).

Discussion

Within the context of annotating a new genome, iLoci provide a quick and convenient solution for leveraging genomes of related model organism to establish baseline expectations about genome composition and organization for the organism of interest. Similarities to genomes of related species across a broad range of measures gives one confidence in the quality of the genome assembly and annotation. By contrast, any stark

differences should point to specific genomic features that warrant additional investigation to distinguish the effects of annotation from real differences in genome biology. Considerable effort has been devoted to making such comparisons as easy as possible: relevant software is freely available as open source code, is engineered with a focus on resource efficiency (enabling it to run easily on laptop or desktop computers), and works with a small number of standard input files. In short, iLoci provide a "common currency" for evaluating new data sets and re-evaluating previously published data sets alike.

Additional applications of iLoci in the annotation and analysis of novel genomes abound. Leveraging iLoci with strong support from expression and homology evidence to train species-specific gene prediction models can yield improvements in subsequent annotation efforts. The longest regions of the genome annotated as intergenic can yield insight into the proliferation of transoposable and other repetitive elements and non-coding RNA genes, or alternatively characteristics of regions where annotation workflows fail to predict genes. The largest regions of high gene density, as represented by miLoci, provide an excellent starting place for investigating the clustering of functionally related genes, whereas miLoci containing two genes are candidates for analysis of tandem gene duplication.

iLoci also facilitate analysis of genome organization at different scales. At the scale of whole chromosomes (or large fractions thereof), iLoci provide a well-defined measure of genome compactness that can be compared across annotations, assemblies, and species. At a slightly smaller scale, iLoci can be leveraged to investigate large-scale changes in genome organization along the length of a chromosome, and possible interpretations in terms of transposon activity and other dynamic mechanisms of genome expansion and contraction. At the scale of individual genes, iLoci capture local aspects of genome organization, furnishing insight into gene spacing and orientation for specific genes of interest

TODO: simulation of genome dynamics.

Conclusions

Parsing annotated genome sequences into iLoci and then using these iLoci as a new coordinate system provides a robust and reproducible framework for investigating a variety of questions about genome content, architecture, and evolution. iLocus annotation might include contextual information for gene models in the form of up- and down-stream regulatory sequences. iLoci containing overlapping gene models can easily be identified for scrutiny seeking to distinguish gene model prediction errors from true compact gene organization that would likely be missed if analysis were performed at the level of individual genes. iLoci also provide stability across different versions of an annotated genome assembly, preserving gene models or intergenic regions for which local genomic context remained invariant to assembly and annotation updates. Finally, iLoci provide a way to break down the entire genome into distinct blocks that can be filtered based on their composition, gene content, conservation, or a variety of other characteristics of interest, thus providing finely tuned data sets for analyses or training and testing of predictive models.

Figures

Figure 4.1

Designation of iLocus types shown in green, with classification logic described in blue.

Figure 4.2

Parsing an annotated genome sequence into iLoci. The letters A to J indicate 10 adjacent iLoci on the genomic sequence (central horizontal line), separated by the long vertical bars. Gene annotations are shown underneath the genome sequence. Exons are schematized by horizontal lines, introns by the triangular thin lines below. Arrows indicate transcriptional direction. iLoci A, C, E, G, and J are without gene annotation, with A and J representing potentially incomplete genomic fragments (fiLoci), and C, E, and G representing intergenic regions (iiLoci). siLoci contain a single gene annotation each and include genes with a single annotated transcript (B, H, and I) as well as genes with alternative transcripts (D). ciLocus F contains three distinct, but overlapping genes. The boundaries of the gene-containing iLoci (giLoci) are derived from the annotation ends, extended in each direction by δ . An exception occurs between iLoci H and I, where the extension would result in an iiLocus shorter than δ : in this case, the bordering iLoci (H and I) are extended towards each other to fill the entire space.

Figure 4.3

Aggregate lengths of n adjacent iiLoci for 3 species (Caenorhabditis elegans, Medicago truncatula, and Mus musculus). The value of n at which the distribution of aggregate iiLocus lengths is normal is reflective of the range at which local variations in gene spacing are evened out.

Figure 4.4

Theoretical values of σ (the proportion of genome sequence occupied by miLoci) plotted as a function of ϕ (the proportion of giLoci merged into miLoci) at different values of ρ (the ratio of iiLocus length to giLocus length).

Figure 4.5

The genomic compactness of ten model organisms, as measured on long (≥ 1 Mb) chromosome or scaffold sequences.

Figure 4.6

Comparison of genome compactness as measured on genomic sequences as annotated (solid squares) versus the same sequences with a random arrangement of genes (hollow circles). Each point represents the average (centroid) of (ϕ, σ) values computed on all long chromosome and scaffold sequences in the corresponding assembly, as shown in Figure 4.5.

Figure 4.7

A breakdown of iLoci from two species by stability and classification.

Figure 4.8

Scatter plot showing the length and nucleotide composition of iLoci from *Arabidopsis* thaliana (top) and *Apis mellifera* (bottom). iLoci that are stable between two assembly/annotation versions are represented by red marks, while unstable iLoci are represented by blue and green marks. siLoci are shown on the left, and iiLoci are shown on the right.

Figure 4.9

Counts and genomic space occupied of iLoci from 9 species of green algae, categorized according to gene content and homology status.

Figure 4.10

Plot of length and nucleotide composition for piLoci from the *Chlamydomonas reinhardtii* genome. Blue point represent *highly conserved piLoci*, green represent *conserved piLoci*, and red represent *orphan piLoci*.

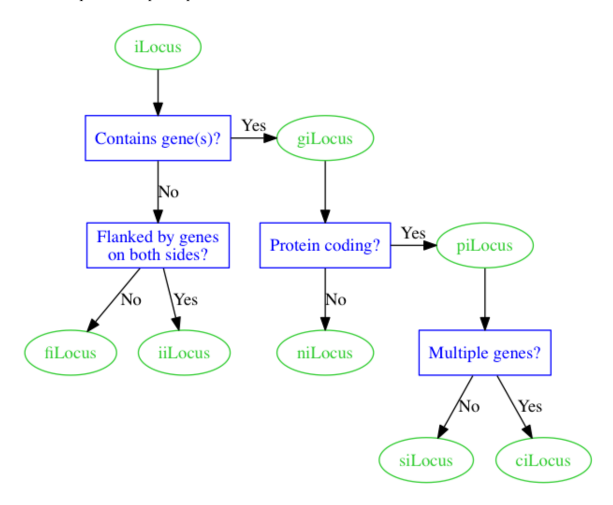


Figure 4.1

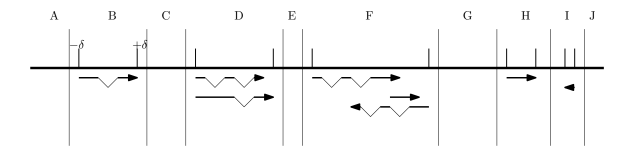


Figure 4.2

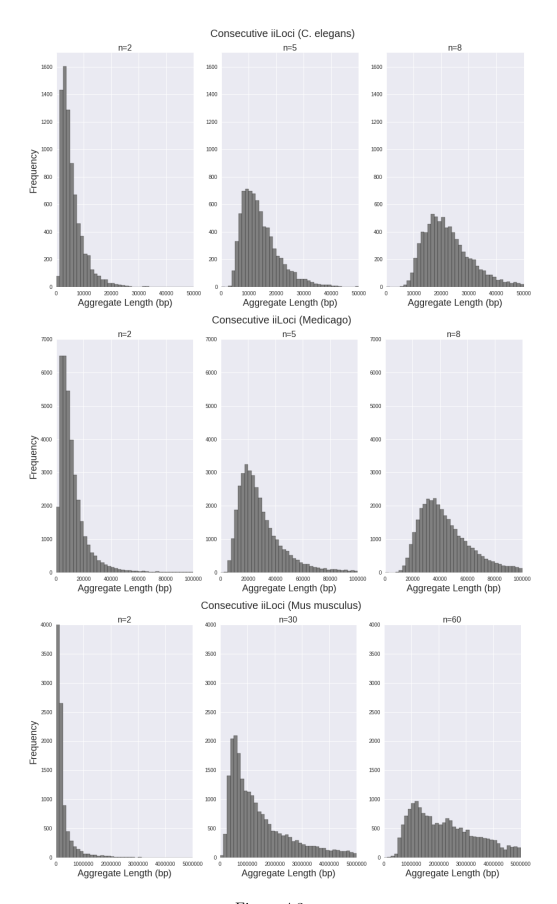


Figure 4.3

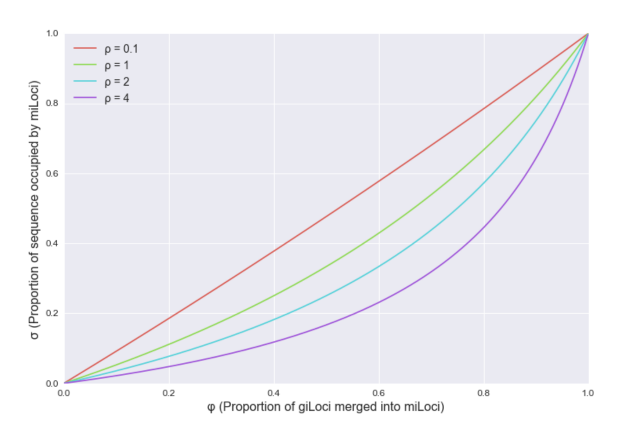


Figure 4.4

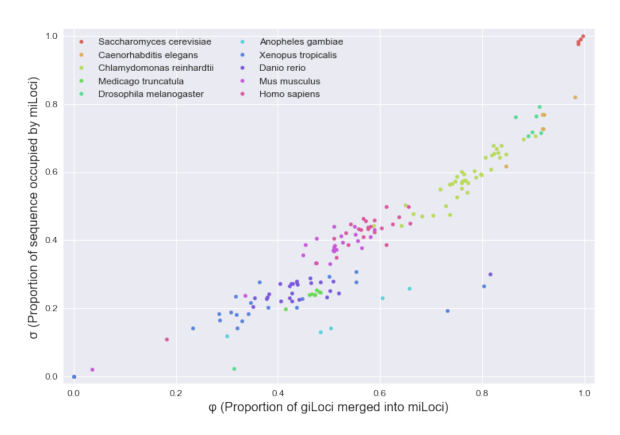


Figure 4.5

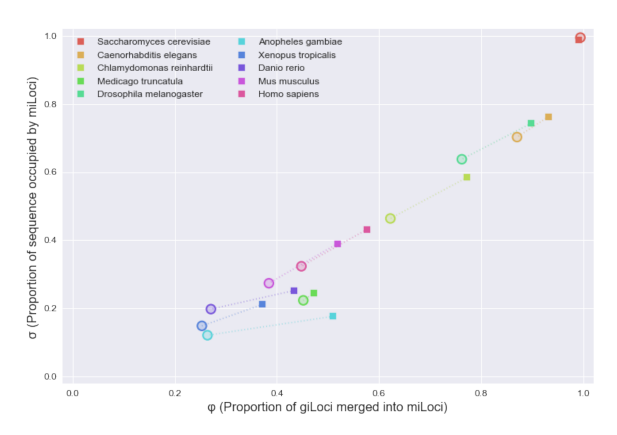
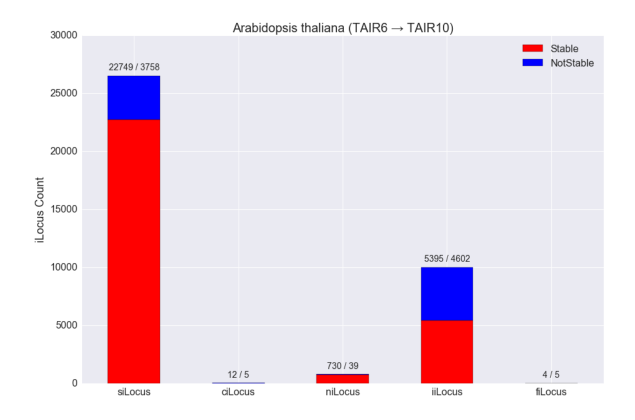


Figure 4.6



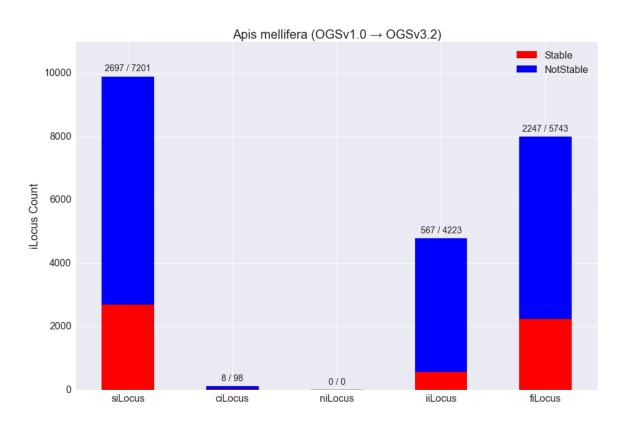


Figure 4.7

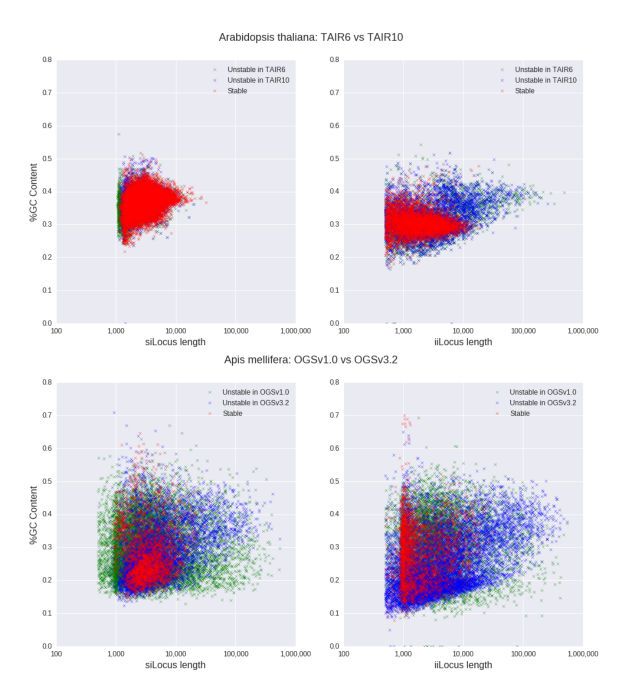
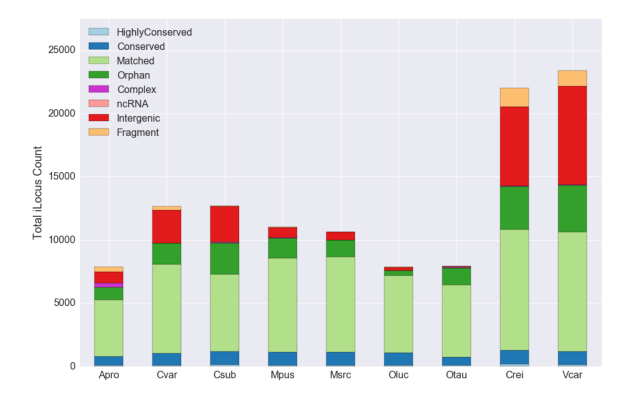


Figure 4.8



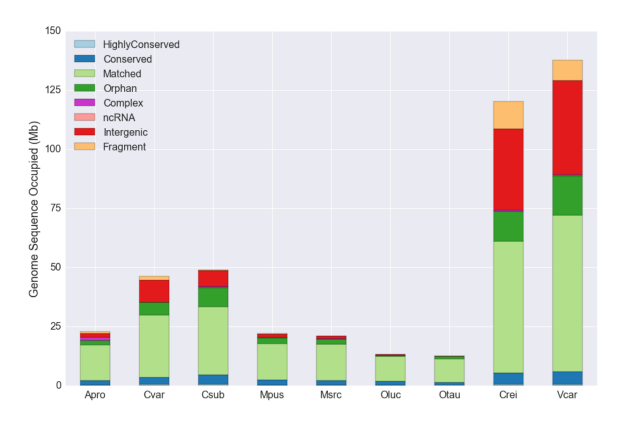


Figure 4.9

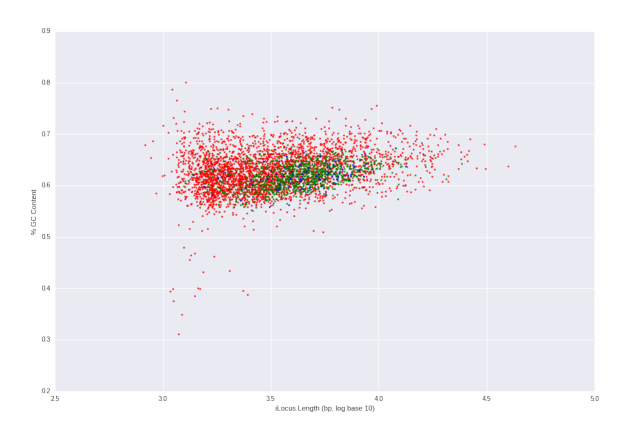


Figure 4.10

CHAPTER 5. CONCLUSIONS

A draft genome assembly and corresponding annotation are important data resources for any genome project, and provide the coordinate systems and catalogs of functional elements upon which genomics studies rely. At the same time, it is widely acknowledged that genome assembly and annotation are complex and difficult with current nucleotide sequencing technologies, requiring close collaboration between domain experts and bioinformaticians. Single-molecule long-read sequencing platforms promise exciting improvements in genome assembly, and within a few years could facilitate a much more direct path to high-quality reference genome sequences. As the cost and effort corresponding to genome-scale sequencing continues to drop, the demands imposed on domain experts to interpret genome-scale data will only continue to increase. The focus of this dissertation has been the development of tools and methodologies that enable reproducible analysis of annotated genomes, as motivated by challenges encountered in assembling, annotating, and analyzing the genome of the paper wasp *Polistes dominula*.

Polistes is an important model for the study of social behavior and its evolution. The genome of Polistes is similar to other social insects in many characteristic features, while showing a slight reduction in others, such as intron length and genome size [119]. The nucleotide composition of Polistes dominula is the most AT-rich of any social insect studied, raising compelling questions about the molecular evolution of the genome and possible contributions of biased repair/mutation mechanisms or historically elevated levels of DNA methylation and deamination. Analysis of gene expression in queen and worker wasps identified 367 caste differentially expressed loci, with functions enriched

for metabolism and neurotransmitter activity, but did not discover any caste bias in alternate splicing. Most striking in terms of genome content is the conspicious absence of the DNA methyltransferase Dnmt3 (recently and independently confirmed as missing in another Polistes genome [42]) and essentially zero DNA methylation genome-wide. Previously thought to be a hallmark signature of social behavior [80], Polistes is the first social insect known to lack genome-wide DNA methylation a complete complement of DNA methylation enzymes.

While intriguing in the context of insect sociogenomics, the challenges I encountered in this project are common challenges in the wider context of genome biology. I therefore sought to create tools that could be applied generally to address common questions about genome composition and organization. Foremost, researchers need to be able to scan newly produced genomes for genes of interest—for example, established "social behavior" genes or DNA methylation related enzymes, as in *Polistes*. Researchers must also be able to compare different sources of annotation for the genome, and assess these in the context of available expression data and other evidence. There must be a way for researchers to refine annotations at loci of interest as needed, and to track changes to the annotation over time, providing complete providence for each data point included in an analysis. And finally, researchers need to be able to select subsets of data with which to reliably describe characteristics of the genome, such as (for example) intron length or codon usage.

The framework provided by *interval loci* (*iLoci*) and related tools is designed to provide precise solutions to these challenges. iLoci furnish a "parts list" of the genome, a representation of the genome that is granular, complete, and that captures the genomic context of each genic or intergenic region. The ParsEval tool [129] identifies the extent to which alternative sources of annotation for the genome differ, and in concert with community annotation tools (such as those provided by xGDBvm [130]) facilitates targeted evaluation and refinement of genes of interest. iLocus statistics produced by GenHub

programs [131] enable filtering of iLoci based on their length, nucleotide composition, and gene content. A prototype of a related tool, GeneAnnoLogy, implements a version control system to track changes in annotations over time, and could be extended with more flexible and comprehensive iLocus filtering mechanisms.

The tools described in this dissertation are designed to calculate and present characteristics of genome composition quickly and easily for a new genome, and to facilitate uniform re-analysis of related data sets for comparison. Accordingly, care was taken to ensure that the tools support standard data formats, and to reduce the friction involved in installing and executing the software. These practical concerns, while largely orthogonal to the science, can nevertheless make "quick and easy" processing and—ultimately—interpretation of genome data prohibitively difficult. Just as improvements in DNA sequencing technology have democratized genome sequencing, making these tools accessible and compatible with well-defined standard data formats puts the power of genome analysis directly in the hands of domain experts. Rapid assessment of new genome assemblies and annotations in the context of related genomes will help researchers distinguish between genomic features that are broadly conserved and those that appear to be unique, warranting additional attention and scrutiny.

In addition to questions of genome composition, iLoci allow researchers to probe questions related to genome organization. In particular, iLoci provide insight into the spacing and orientation of genes at the whole-genome scale, and provide well-defined measures of genome compactness that are remarkably consistent between different genome sequences within the same species, and between different species within the same clade. While these insights are investigated primarily in a descriptive context, exciting possibilities await in exploring and applying these insights in modeling and potentially even genetic engineering. Are there certain features of genome organization associated with traits of interest, and can these be genetically engineered in an organism? Or, by simulating transposon activity, gene duplication, genome rearrangement, mutation, and potentially

even whole genome duplication, can we produce artificial genomes that bear signatures of large-scale genome organization observed in real genomes?

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