Genome analysis

A computational genomics pipeline for prokaryotic sequencing projects

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ABSTRACT

Motivation: New sequencing technologies have accelerated research on prokaryotic genomes and have made genome sequencing operations outside major genome sequencing centers routine. However, no off-the-shelf solution exists for the combined assembly, gene prediction, genome annotation, and data presentation necessary to interpret sequencing data. The resulting requirement to invest significant resources into custom informatics support for genome sequencing projects remains a major impediment to the accessibility of high-throughput sequence data.

Results: We present a self-contained, automated high-throughput open source genome sequencing and computational genomics pipeline suitable for prokaryotic sequencing projects. The pipeline has been used at the Georgia Institute of Technology and the Centers for Disease Control and Prevention for the analysis of Neisseria meningitidis and Bordetella bronchiseptica genomes. The pipeline is capable of enhanced or manually assisted reference-based assembly using multiple assemblers and modes; gene predictor combining; and functional annotation of genes and gene products. Because every component of the pipeline is executed on a local machine with no need to access resources over the Internet, the pipeline is suitable for projects of a sensitive nature. Annotation of virulence-related features makes the pipeline particularly useful for projects working with pathogenic prokaryotes.

Availability and Implementation: The pipeline is licensed under the open-source GNU General Public License and available at the Georgia Tech Neisseria Base (http://nbase.biology.gatech.edu/). The pipeline is implemented with a combination of Perl, Bourne Shell, and MySQL and is compatible with Linux and other Unix systems.

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Supplementary information: See http://nbase.biology.gatech.edu

1 INTRODUCTION

Genome sequencing projects, pioneered in the 1990s (Fleischmann, et al., 1995), require large-scale computational support in order to make their data accessible for use and interpretation by biologists. Large sequencing centers have traditionally employed or collaborated with teams of software engineers and computational biologists to develop the software and algorithms for sequencing hardware interfaces, enterprise data storage, sequence assembly and finishing, genome feature prediction and annotation, database mining, comparative analysis, and database user interface development. While many of the components developed by these teams are now available online under open-access terms, the development of new, high-throughput sequencing technologies has necessitated updates to these tools and development of even more sophisticated algorithms to address the challenges raised by the new data. These new technologies – 454 pyrosequencing (Margulies, et al., 2005), ABI SOLiD (Shendure, et al., 2005), and Illumina (Bentley, et al., 2008) – are now collectively referred to as second generation sequencing technologies. Similar updates will be needed as the third generation of sequencing technologies, such as Pacific Biosciences’ SMRT sequencing (Eid, et al., 2009), enter production use. New and improved tools released for these technologies on a monthly basis include assemblers, mapping algorithms, base calling and error correction tools, and a multitude of other programs. Because of this fast pace of development, few experts are able to keep up with the state of the art in the field of computational genomics. Accordingly, the rate limiting step in genome sequencing projects is no longer the experimental characterization of the data but rather the availability of experts and resources for computational analysis.

At the same time, the increased affordability of these new sequencing machines has spawned a new generation of users who were previously unable to perform their own genome sequencing, and thus collaborated with large sequencing centers for genome sequencing and subsequent computational analysis. While these users are now able to experimentally characterize genomes in...
house, they often find themselves struggling to take full advantage of the resulting data and to make it useful to the scientific community since the informatics support for their genome projects is not sufficient.

Several large sequencing consortia (Aziz, et al., 2008; Markowitz, et al., 2009; Seshadri, et al., 2007) have produced comprehensive, centralized web-based portals for the analysis of genomic and metagenomic data. While extremely useful for many types of projects and collaborations, these solutions inherently result in a loss of data processing flexibility compared to locally installed resources and may be unsuitable for projects dealing with sensitive data. Recently, another group (Stewart, et al., 2009) has published DIYA, a software package for gene prediction and annotation in bacterial genomes with a modularized, open source microbial genome processing pipeline. However, DIYA does not include a genome assembly component, and does not provide for the combination of complementary algorithms for genome analysis.

To address the outstanding challenges for local computational genomics support, we have developed a state of the art, self-contained, automated high-throughput open source software pipeline for computational genomics in support of prokaryotic sequencing projects. To ensure the relevance of our pipeline, we checked the latest developments in computational genomics software for all stages of the pipeline, such as new versions of assembly and gene prediction programs and comparative surveys, and selected what we deemed to be the most suitable software packages. The pipeline is self-contained; that is, we used locally installable versions of all third-party tools instead of web-based services provided by many groups. We chose to do so for three reasons: first, because some of the applications we envision for this pipeline are of sensitive nature; second, to enhance robustness to external changes (e.g., online API changes or website address changes); and third, to improve the ability of developers to customize and derive from our pipeline. The pipeline is also automated and high-throughput: all components are organized in a hierarchical set of readily modifiable scripts, and the use of safe programming practices ensures that multiple copies of the pipeline can be run in parallel, taking advantage of multiple processors where possible.

Importantly, by using and combining the outputs of competitive, complementary algorithms for multiple stages of genome analysis, our pipeline allows for substantial improvement upon single-program solutions. The use of multiple algorithms also provides a way to improve robustness and conduct more comprehensive quality control when the output of one program is significantly different from that of another.

Computational support provided to prokaryotic genome projects by our pipeline can be subdivided into three stages: first, sequencing and assembly; second, feature prediction; and third, functional annotation. For the assembly stage, we developed a custom protocol specific to 454 pyrosequencing data, which resulted in a significant improvement to assembly quality of our test data compared to the baseline assembler bundled by the manufacturer. Other assemblers can be plugged in if necessary, and data from other sequencing technologies such as ABI SOLiD, Illumina and Sanger capillary-based machines can be used. For the prediction stage, we again included a custom combination of feature prediction methods for protein-coding genes, RNA genes, operon and promoter regions, which improves upon the individual constituent methods.

The annotation stage includes several types of protein functional prediction algorithms. We also developed components for comparative analysis, interpretation and presentation (a web-based genome browser), which can be used downstream of our pipeline.

We have tested the pipeline on the bacterium Neisseria meningitidis, which is a human commensal of the nasopharynx and which can sometimes cause meningitis or septicaemia (Rosenstein, et al., 2001). When N. meningitidis does cause disease, it can be devastating with an approximately 10% fatality rate and 15% sequelae rate. N. meningitidis is a highly competent organism with a high recombination rate, and large chromosomal changes are common (Jolley, et al., 2005; Schoen, et al., 2008). This complicates computational genome analysis and makes N. meningitidis an appropriately challenging test for our pipeline. To demonstrate the general applicability of the pipeline, we have also tested it on a different pathogen, Bordetella bronchiseptica. B. bronchiseptica is a Gram-negative bacterium that can cause bronchitis in humans, although it is more commonly found in smaller mammals (Parkhill, et al., 2003). Much like Neisseria, Bordetella has extensive plasticity, likely due to the large number of repeat elements (Gerlach, et al., 2001). Here, we analyze the first two complete genome sequences of B. bronchiseptica strains isolated from human hosts.

The rest of this paper is organized as follows. The System and Methods section describes the genomes which we used to test our pipeline, overall organization of the pipeline, and details of the algorithms used to perform tasks in the pipeline. In the Discussion section, we discuss the objectives of our work on the pipeline and how these relate to larger developments in computational biology for next-generation sequencing.

2 SYSTEM AND METHODS

2.1 Genome test data

N. meningitidis genomes were characterized via 454 pyrosequencing (Margulies, et al., 2005) using either a half or one quarter plate runs on the Roche 454 GS-20 or GS Titanium instrument (Table 1). For each genome, a random shotgun library was produced using Roche protocols for nebulization, end-polishing, adaptor ligation, nick repair and single-stranded library formation. Following emulsion PCR, DNA bound beads were isolated and sequenced using long read (LR) sequencing kits. The number of reads produced in the experiments ranged from 200,000 to 600,000, and the average read lengths were between 100 and 330 bases. These data yielded 47.6-94.3 million bases per genome amounting to 20-40x coverage for the approx. 2.1 megabase N. meningitidis genomes. After read trimming and re-filtering to recover short quality reads, the data were passed to the first stage of the pipeline – genome assembly.

2.2 Pipeline organization

The analytical pipeline consists of three integrated subsystems: genome assembly, feature prediction and functional annotation. Each subsystem consists of a top-level execution script managing the input, output, format conversion, and combination of results for a number of distinct software components. A hierarchy of scripts and external programs then performs the tasks required to complete each stage of analysis (Figure 1).
2.3 Assembly

Genome assembly was performed by evaluating multiple configurations of assemblers including the standard 454 assembler, Newbler (version 2.3), as well as the Celera Assembler (Miller et al., 2008), the Phrap assembler (http://www.phrap.org/) and the AMOScmp mapped assembler (Pop et al., 2004). Several other assemblers were evaluated but ultimately excluded from the pipeline due to use limitations; for instance, the ALLPATHS 2 assembler (MacCallum et al., 2009) required paired-end reads to operate; our evaluation data contained no paired-end reads, and such a requirement unnecessarily constrains the user’s options. The widely used Velvet assembler (Zerbino and Birney, 2008) was originally developed as de novo assembler for Illumina sequencing technology, but its capability has been extended to accommodate 454 data as well. However, we were unable to configure the Velvet assembler to produce a usable assembly or take advantage of reference genomes using 454 data alone.

Evaluation of the results indicated that mapped assemblies of N. meningitidis genomes using previously finished strains were of superior quality to de novo assemblies. Using the most appropriate reference strains, it was found that Newbler and AMOScmp complement each other’s performance in the assembly stage, with
Newbler being able to join some contigs AMOScmp left gapped and vice versa. As a result, we decided to use a combination of these two assemblers’ outputs for the final assembly. Then, the Minimus assembler (Sommer, et al., 2007) from the AMOS package, a simple assembler for short genomes, was used to combine the constituent assemblies.

We also evaluated alternative base calling algorithms for 454 pyrosequencing data (Quinlan, et al., 2008) but detected no improvement. Over the course of our project, accuracy of base calling in the Newbler assembler was reported to be significantly improved. We used the latest version of the assembler available at publication time (2.3).

An optional component of the pipeline was created for frameshift detection using FSFind (Kislyuk, et al., 2009). Frameshifts in protein-coding sequences are a known result of pyrosequencing errors caused by undercalls and overcalls in homopolymer runs (Kuo and Grigoriev, 2009). Briefly, this package creates a GeneMark model of the genome, makes gene predictions, and then scans the genome for possible frameshift positions on the basis of ORF configuration and coding potential. Once the possible frameshift sites are identified, a putative translation of the protein possibly encoded by the broken gene is compared against a protein database (SwissProt by default). The predicted frameshift site is also scanned for adjacent homopolymers. A heuristic set of confidence score cutoffs is then used to provide a set of frameshift predictions while minimizing the false positive rate. The predicted frameshift sites can then be verified experimentally or corrected speculatively. The user can inspect the dataset to decide whether locations predicted to contain frameshifts break gene models, and patch the sequences to fix up these positions. The prediction stage can then be re-run to correct the gene predictions. While further experimental analysis to address such errors is desirable (e.g., targeted PCR of predicted error locations or a recently popular choice of combining sequencing technologies such as 454 and Illumina), it incurs extra costs which we aim to avoid.

Unfinished assemblies produced in this stage contained 90-300 contigs each. No paired-end libraries or runs were available for the strains analyzed, and therefore scaffolding of the contigs was a challenge. Manual examination of the assemblies using the MAUVE (Darling, et al., 2004) multiple whole-genome alignment and visualization package revealed numerous locations where contigs could be scaffolded with a small gap or minimal overlap (Figure 2). As an optional step, we produced a table of such positions and a script which would scaffold contigs joined by the gap.

Table 2. Summary of assembler performance. Data for each strain are presented in rows. Statistics from standalone assemblers (Newbler and AMOScmp) are presented together with results of the combining protocol (default output of the pipeline) and an optional, manually assisted predictive gap closure protocol.

<table>
<thead>
<tr>
<th>Strain ID</th>
<th>Newbler statistics</th>
<th>AMOScmp statistics</th>
<th>Automatic combined assembly</th>
<th>Manual combined assembly</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Contigs &gt; 500 nt, total size</td>
<td>Contigs &gt; 500 nt, N50, Longest contig</td>
<td>Contigs &gt; 500 nt, total size</td>
<td>Contigs &gt; 500 nt, N50, Longest contig</td>
</tr>
<tr>
<td>NM13220</td>
<td>172</td>
<td>22K</td>
<td>202</td>
<td>21K</td>
</tr>
<tr>
<td>NM10699</td>
<td>102</td>
<td>52K</td>
<td>116</td>
<td>43K</td>
</tr>
<tr>
<td>NM15141</td>
<td>2.06M</td>
<td>143K</td>
<td>190</td>
<td>1.13K</td>
</tr>
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<td>33K</td>
<td>190</td>
<td>22K</td>
</tr>
<tr>
<td>NM18575</td>
<td>2.09M</td>
<td>38K</td>
<td>147</td>
<td>29K</td>
</tr>
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<td>NM1578</td>
<td>2.09M</td>
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<td>88K</td>
</tr>
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<td>NM15293</td>
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<td>42K</td>
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<td>110</td>
<td>42K</td>
</tr>
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<td>BBF579</td>
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<td>136K</td>
<td>132K</td>
<td>1.2M</td>
</tr>
</tbody>
</table>

(a) N50 is a standard quality metric for genome assemblies that summarizes the length distribution of contigs. It represents the size N such that 50% of the genome is contained in contigs of size N or greater. Greater N50 values indicate higher quality assemblies. (b) No improvement was detected from the combined assembly in strain BBF579, and the original Newbler assembly was automatically selected. (c) The manual combined assembly protocol was not performed for these projects.
Then, a manual gap joining stage used the layout of the contigs according to their aligned positions on the reference using the AMOS package and manual examination of each gap, adjacent contig alignments and reference annotation in the MAUVE visualization tool. Although there is a possibility that rearrangements exist in those gaps as mapped to the closest reference genome, joining was only done after manual examination on a case-by-case basis in positions of high homology and full consensus between four of the reference strains, to minimize this possibility. While we provide the scripts and data format definitions necessary to complete this stage of the pipeline, it involves manual processing of the assembly and is therefore optional. This component is similar in function to Mauve Contig Mover (Rissman, et al., 2009) but expands upon it in several ways. An option is provided in the pipeline to use Mauve Contig Mover.

The manually assisted genome assembly procedure resulted in an order-of-magnitude decrease in the number of gaps in comparison to the Newbler assembler (which in turn performed the best out of all standalone assemblers evaluated). In addition, the fully automated assembly metrics (N50 and contig count at equal minimal size) are an approximately 20-50% improvement upon baseline Newbler performance (Table 2).

The contigs in the assembly stage output were named according to the following format: prefix_contig#, where the prefix represents a unique strain identifier and # represents the zero-padded sequential number indicating the contig’s predicted order on the chromosome. For example, the 25th contig for the N. meningitidis strain M13220 assembly would be named as CDC_NME_M13220_025. The prefix used in the pipeline is configurable by the user with a command line option.

2.4 Feature prediction

Feature prediction was performed in the genome using a suite of several programs. To predict genes, we used a combination of de novo and comparative methods. The Glimmer (Delcher, et al., 1999) and GeneMark (Besemer, et al., 2001) microbial gene predictors were used for de novo prediction, and BLASTp alignment (Altschul, et al., 1997) of putative proteins was used for comparative prediction. Self-training procedures were followed for both de novo predictors, and the results, while highly concordant, were different enough (Table 3) to justify the inclusion of both algorithms. BLASTp alignment of all open reading frames (ORFs) at least 90 nt long was performed using the Swiss-Prot protein database (Boeckmann, et al., 2003).

The results of these three methods were combined together using a combiner strategy outlined in Figure 3. In this strategy, we first check that at least half of the predictors report a gene in a given ORF – in our configuration, 2 of the 3 predictors. Then the Met (putative translation start) codon closest to the beginning of the BLAST alignment is found and declared to be the gene start predicted by BLAST. We then find the gene start coordinate reported by the majority of the three predictors and report the resulting gene prediction. If no majority exists, we select the most upstream gene start predicted.

![Fig. 3. Schematics of combining strategy for prediction stage. BLAST alignment start, which may not coincide exactly with a start codon, is pinned to the closest start codon. Then, a consensus or most upstream start is selected.](http://bioinformatics.oxfordjournals.org)

<table>
<thead>
<tr>
<th>Strain ID</th>
<th>Gene predictions by GeneMark</th>
<th>Gene predictions by Glimmer3</th>
<th>Gene predictions by BLAST</th>
<th>ORFs with full consensus</th>
<th>ORFs with partial consensus</th>
<th>Total gene predictions reported</th>
<th>tRNAs predicted by tRNAscan-SE</th>
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<td>NM13220</td>
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<td>NM10699</td>
<td>2366</td>
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<td>1317</td>
<td>1284</td>
<td>826</td>
<td>2110</td>
<td>51</td>
</tr>
<tr>
<td>NM15141</td>
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<td>2578</td>
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<td>1343</td>
<td>841</td>
<td>2184</td>
<td>57</td>
</tr>
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<td>1341</td>
<td>1308</td>
<td>802</td>
<td>2110</td>
<td>51</td>
</tr>
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<td>2927</td>
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<td>1448</td>
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<td>63</td>
</tr>
<tr>
<td>NM5178</td>
<td>2377</td>
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<td>1315</td>
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<td>816</td>
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<td>52</td>
</tr>
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<td>1285</td>
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<td>802</td>
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<td>51</td>
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<td>4793</td>
<td>4793</td>
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<td>2732</td>
<td>2067</td>
<td>4799</td>
<td>48</td>
</tr>
<tr>
<td>BBF579</td>
<td>4649</td>
<td>4646</td>
<td>2652</td>
<td>2635</td>
<td>2021</td>
<td>4656</td>
<td>48</td>
</tr>
</tbody>
</table>

(a) Number of ORFs with protein-coding gene predictions where all 3 predictors agreed exactly or with a slight difference in the predicted start site. (b) ORFs where only 2 of the 3 predictors made a prediction. (c) Total protein-coding gene predictions reported by the pipeline.

In addition to protein-coding gene prediction, ribosomal genes were predicted using alignment to a reference database of ribosomal operons, and tRNA genes were predicted using the tRNAscan-SE package (Lowe and Eddy, 1997). The results are summarized in Table 3.

Results of the feature prediction stage are saved in a multi-extent GenBank formatted file. Features were named according to the
following convention: contig-name_feature-id, where contig-name is as described earlier, and feature-id is a sequential zero-padded number unique to the feature across all contigs. For example, a gene with feature ID 1293 on contig 25 might have the name CDC_NME_M13320_025_1293.

To validate the overall accuracy of the gene prediction stage of the pipeline, we ran our gene prediction tools on the genome of *Escherichia coli* K12, one of the best-annotated bacterial genomes (analysis described in the Supplementary Text document). Our pipeline was able to detect 95.7% of the annotated *E. coli* K12 protein-coding genes, and exactly predict starts in 85.5% of those. 50% of the *E. coli* predictions that report incorrect start codons report starts within 35 nt of the true start, and all reported starts are within 200 nt of the true start.

### 2.5 Functional annotation

Functional annotation of genome features was also performed using a combination of tools. Annotation of protein coding genes was based on an integrated platform that makes use of six distinct annotation tools, four of which employ intrinsic sequence characteristics for annotation and two that use extrinsic homology-based approaches to compare sequences against databases of sequences and structures with known functions. Information on Gene Ontology (GO) terms, domain architecture and identity, subcellular localization, signal peptides, transmembrane helices and lipoprotein motifs is provided for each protein coding gene (Figure 4). BLASTp alignment of predicted proteins was performed against the UniProt database (Uniprot, 2009). Homology-based searches were also made across thirteen sequence and protein domain databases with the InterProScan suite (Mulder and Apweiler, 2007). Parsing of the results was carried out against the corresponding InterPro database. The pipeline also stores the top five hits for each gene against the NCBI non-redundant protein database, to provide potentially useful information. All homology searches were run locally. Signal peptides were annotated using the SignalP package (Bendtsen, et al., 2004) and transmembrane domains were annotated with the TMHMM package (Krogh, et al., 2001). State of the art in subcellular localization algorithms was examined to ensure the best performance given our operational requirements. Insertion sequences (transposons) and proteins reported as virulence factors by VFDB (Chen, et al., 2005; Yang, et al., 2008) were also annotated. These annotations of virulence-related features make the pipeline particularly useful for projects working with pathogenic prokaryotes. Results of this analysis are summarized in Table 4.

After the functional annotations were determined, a naming scheme was employed for each locus to conform to standard annotation terminology. Specific gene names were assigned according to homology-based results. For genes that had a UniProt result with a best hit at greater than 91% amino acid sequence identity and an e-value less than 1e-9, the gene assumed the best hit’s name. If the best hit had the keyword “hypothetical,” then we used a domain name from the homology database.

<table>
<thead>
<tr>
<th>Strain ID</th>
<th>Total number of CDS</th>
<th>Signal peptides</th>
<th>Transmembrane helices</th>
<th>Conserved hypothetical proteins</th>
<th>Putative uncharacterized proteins</th>
<th>Functional assignment inferred from homology</th>
<th>Virulence factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>NM10699</td>
<td>2110</td>
<td>310 (14.7%)</td>
<td>180 (8.5%)</td>
<td>5 (0.2%)</td>
<td>652 (30.9%)</td>
<td>577 (27.3%)</td>
<td>45 (2.1%)</td>
</tr>
<tr>
<td>NM15141</td>
<td>2184</td>
<td>317 (14.5%)</td>
<td>173 (7.9%)</td>
<td>16 (0.7%)</td>
<td>590 (27.0%)</td>
<td>583 (26.7%)</td>
<td>50 (2.3%)</td>
</tr>
<tr>
<td>NM9261</td>
<td>2063</td>
<td>303 (14.4%)</td>
<td>166 (7.9%)</td>
<td>13 (0.6%)</td>
<td>591 (28.0%)</td>
<td>558 (26.4%)</td>
<td>37 (1.8%)</td>
</tr>
<tr>
<td>NM18575</td>
<td>2471</td>
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<td>193 (7.8%)</td>
<td>13 (0.5%)</td>
<td>725 (29.3%)</td>
<td>668 (27.0%)</td>
<td>48 (1.9%)</td>
</tr>
<tr>
<td>NM5178</td>
<td>2097</td>
<td>298 (14.2%)</td>
<td>177 (8.4%)</td>
<td>3 (0.1%)</td>
<td>646 (30.8%)</td>
<td>572 (27.3%)</td>
<td>45 (2.1%)</td>
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<tr>
<td>NM15293</td>
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<td>168 (8.1%)</td>
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<td>613 (29.7%)</td>
<td>567 (27.5%)</td>
<td>47 (2.3%)</td>
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<tr>
<td>BBE001</td>
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<td>368 (7.7%)</td>
<td>9 (0.2%)</td>
<td>807 (16.8%)</td>
<td>1184 (24.7%)</td>
<td>54 (1.1%)</td>
</tr>
<tr>
<td>BBF579</td>
<td>4656</td>
<td>934 (20.1%)</td>
<td>339 (7.3%)</td>
<td>9 (0.2%)</td>
<td>739 (15.9%)</td>
<td>1171 (25.2%)</td>
<td>45 (1.0%)</td>
</tr>
</tbody>
</table>

(a) Total putative protein-coding sequences analyzed. (b) As predicted by SignalP (Bendtsen, et al., 2004); percentage of total CDS indicated in parentheses. (c) As predicted by TMHMM (Krogh, et al., 2001). (d) As predicted by BLASTp alignment against VFDB (Chen, et al., 2005; Yang, et al., 2008); http://www.mgc.ac.cn/VFs/
name from InterPro to name the gene. For example, if a gene was given the name “hypothetical” from Uniprot and a domain name of “transferase” from InterPro, then the final name was “hypothetical transferase protein.” Therefore most genes that were given “hypothetical” or “putative” prefixes could then be given a more comprehensive name based on further information such as domain names or protein functions. Genes with unknown functions found across many genomes were given the name “conserved hypothetical protein,” and all other putative genes with unknown functions were given the name “putative uncharacterized protein.”

2.6 Availability
The pipeline software package is available at our website (http://nbase.biology.gatech.edu). The package contains detailed instructions and scripts for installation of the pipeline and all external programs, documentation on usage of the pipeline and its organization. Components which require large biological databases automatically download local copies of those databases upon installation.

All of the N. meningitidis genomes reported here, along with custom annotations and tools for searching and comparative sequence analysis, are available for researchers online at our genome browser database (http://nbase.biology.gatech.edu).

3 DISCUSSION

3.1 Genome biology of N. meningitidis and B. bronchiseptica
We have used the pathogen N. meningitidis for the majority of developmental and production testing of our pipeline. Although N. meningitidis gains no fitness advantage from virulence, it occasionally leaves its commensal state and causes devastating disease (Meyers, et al., 2003). Several recent studies have used whole-genome analysis to determine the basis of virulence in this species but none have been conclusive (Hotopp, et al., 2006; Perrin, et al., 2002; Schoen, et al., 2008). With the recent advent of next-generation sequencing and the application of an analytical pipeline, such as presented here, this problem and other problems like it can be addressed in individual laboratories on a genome-wide scale. Here, we briefly speculate on a few of the implications of our findings for the genome biology of N. meningitidis to underscore the potential utility of our pipeline.

Whole genome analysis of microbes has led to the development of the ‘pan-genome’ concept (Tettelin, et al., 2005). A pan-genome refers to the collection of all genes found within different strains of the same species. An open pan-genome means that the genome of any given strain will contain unique genes not found within the genomes of other known strains of the same species. The extent to which microbial pan-genomes are open is a matter of debate (Lapierre and Gogarten, 2009). Recent studies have suggested that the N. meningitidis pan-genome is essentially open (Schoen, et al., 2008), consistent with the fact that it is known to be a highly competent species (Chen and Dubnau, 2004; Kroll, et al., 1998). We evaluated this hypothesis by finding the number of unique genes in each of the seven strains reported here along with seven previously published strains, using the results of our analytical pipeline. Our findings are consistent with (Schoen, et al., 2008) in the sense that every genome sequence was found to contain at least 43 unique genes not found in any other strain. Thus, the N. meningitidis pan-genome does appear to be open.

N. meningitidis is a human commensal that most often does not cause disease, and avirulent strains of the species are referred to as carriage strains. Results of previous comparative genomic analyses have been taken to suggest that carriage strains represent a distinct evolutionary group that is basal to a group of related virulent strains of N. meningitidis (Schoen, et al., 2008). We tested this hypothesis using the results of our analytical pipeline applied to three carriage strains and eight virulent strains of N. meningitidis. Whole genome sequences were aligned and pairwise distances between genomes, based on nucleotide diversity levels, were compared within and between groups of carriage and virulent strains. We found that average of the pairwise genome sequence distances within (w) the carriage and virulent groups of strains was not significantly different from the average pairwise distances between (b) groups (w=0.074±0.027 b=0.090±0.014, t=0.693, P=0.491). This result is inconsistent with the previously held notion that carriage and virulent strains represent distinct evolutionary groups based on whole genome analysis. However, our findings are consistent with earlier work that found little genetic differentiation between carriage and virulent strains of N. meningitidis (Jolley, et al., 2005).

Currently, there is no unambiguous molecular assay to distinguish B. bronchiseptica from other Bordetella species. One reason the two B. bronchiseptica genomes reported here were characterized was to discover genes unique to the species (i.e. not present in any other Bordetella species) to facilitate the development of a B. bronchiseptica-specific PCR assay. To identify such genes, we performed BLASTn with B. bronchiseptica query genes uncovered by our pipeline against other B. bronchiseptica strain genomes along with four genomes of closely related Bordetella species. We uncovered a total of 223 genes that are present in all B. bronchiseptica strains and absent in all other Bordetella species. To narrow down this set of potential PCR assay targets, we searched for the most conserved B. bronchiseptica-specific genes. As a point of reference, we determined the sodC gene used in the N. meningitidis-specific PCR assay (Kroll, et al., 1998) to be 99.6% identical among all six completely sequenced strains of N. meningitidis. There are 7 B. bronchiseptica-specific genes with ≥99.6% sequence identity; these genes represent a prioritized list of potential PCR assay targets.

3.2 Computational genomics pipeline
We have presented our computational genomics pipeline, a local solution for automated, high-throughput computational support of prokaryotic genome sequencing projects. While the revolution in sequencing technology makes possible the execution of genome projects within individual laboratories, the computational infrastructure to fully realize this possibility does not yet exist. We made a comprehensive effort to put the tools required for this infrastructure into the hands of biologists working with next-generation sequencing data. Our aim in the course of this project was to facilitate decentralized biological discoveries based on affordable whole-genome prokaryotic sequencing, a mode of science termed ‘investigator-initiated genomics’. For example, one project
enabled by the pipeline in our laboratory is a platform for SNP detection and analysis in groups of bacterial genomes.

One of our major goals was to provide full automation of our pipeline’s entire workflow, and this has been achieved. On the other hand, to allow computationally savvy users to realize the power of customizability, a semi-automated process is desirable. We have made an effort to strike a balance between these objectives, and provide a modular, hierarchically organized structure to permit maximum customization when so desired.

The state of the art in prokaryotic computational genomics moves at a formidable pace. The modular organization of our pipeline, along with the emphasis on integration of complementary software tools, allows us to continually update our platform to keep pace with developments in computational genomics. For instance, if a new, better assembler becomes available, we can include its results in the assembly stage with a simple change to the pipeline code.

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