4Pipe4 – A 454 data analysis pipeline for SNP detection in datasets with no reference sequence or strain information

Francisco Pina-Martins\textsuperscript{1,2*}, Bruno M. Vieira\textsuperscript{1,3}, Sofia G. Seabra\textsuperscript{1}, Dora Batista\textsuperscript{1,4} and Octávio S. Paulo\textsuperscript{1}

Abstract

Background: Next-generation sequencing datasets are becoming more frequent, and their use in population studies is becoming widespread. For non-model species, without a reference genome, it is possible from a panel of individuals to identify a set of SNPs that can be used for further population genotyping. However, the lack of a reference genome to which the sequenced data could be compared makes the finding of SNPs more troublesome. Additionally, when the data sources (strains) are not identified (e.g., in datasets of pooled individuals), the problem of finding reliable variation in these datasets can become much more difficult due to the lack of specialized software for this specific task.

Results: Here we describe 4Pipe4, a 454 data analysis pipeline particularly focused on SNP detection when no reference or strain information is available. It uses a command line interface to automatically call other programs, parse their outputs and summarize the results. The variation detection routine is built-in in the program itself. Despite being optimized for SNP mining in 454 EST data, it is flexible enough to automate the analysis of genomic data or even data from other NGS technologies. 4Pipe4 will output several HTML formatted reports with metrics on many of the most common assembly values, as well as on all the variation found. There is also a module available for finding putative SSRs in the analysed datasets.

Conclusions: This program can be especially useful for researchers that have 454 datasets of a panel of pooled individuals and want to discover and characterize SNPs for subsequent individual genotyping with customized genotyping arrays. In comparison with other SNP detection approaches, 4Pipe4 showed the best validation ratio, retrieving a smaller number of SNPs but with a considerably lower false positive rate than other methods. 4Pipe4’s source code is available at https://github.com/StuntsPT/4Pipe4.

Keywords: Genomics, SNP detection, Next generation sequencing, 454, Pipeline, EST, Non model organisms
for designing genotyping arrays to further explore their data. The pipeline can provide very high quality SNPs as well as the flanking region sequence, necessary for the design of customized genotyping arrays, currently the most efficient way to extend SNP genotyping from those found in a panel of samples to a larger set of individuals for population genomic studies [4, 5].

Due to the nature of NGS data, any automated pipeline has to be strict enough as to follow a work-flow but, at the same time, flexible enough to serve the different purposes of each investigator. This is the role that 4Pipe4 intends to take. Although 4Pipe4 is tuned for EST data, it can also be used with genomic data and, to some extent, to help automate the process of gene discovery.

Implementation

4Pipe4 is written in Python 3 and is licensed under the GPLv3. It is written in a modular manner that allows for relatively simple expansion of functionality.

Most of the functions present in 4Pipe4 result from the automation of already existing programs and the integration of their respective outputs. However, the variation detection routines are of original design and are based on three criteria, all of which can be adjusted by the user:

- **Base coverage** – The minimum required coverage (C); the default value is 15;
- **Base variants** – The minimum number of equal base variants required in a position (vmin); the default value is 20 % of the minimum required coverage;
- **Base quality** – The average minimum quality of each of the base variants (Qmin); the default value is 70.

This means that in order to consider a position of the alignment as a putative SNP, the below condition must be verified:

\[
\sum R \geq C \land \sum V2 \geq v_{\text{min}} \land Q_{V1} \geq Q_{\text{min}} \land Q_{V2} \geq Q_{\text{min}}
\]

Where “R” is the number of reads in the considered position, “C” is the minimum coverage as defined by the user; “V1” is the most frequent variant base type in the considered position. “V2” is the second most frequent variant base type in the considered position and “Q” is the quality value.

4Pipe4 uses a configuration file, called “4Pipe4rc” with a simple and self documented syntax for setting variables such as the location of programs, the SNP detection criteria and the parameters that should be passed to the external software. How the program uses this configuration file is explained in detail in the program documentation.

The analysis process is divided in 9 steps, each of which can be excluded from the run by issuing the appropriate arguments at run time. In step 1, 4Pipe4 takes an SFF file and, if all the steps are run, step 9 outputs a series of HTML formatted reports, compressed in 7zip. Steps 7 (Gene Ontology) and 8 (SSR detection) are considered optional since they are not required for the SNP detection routines. 4Pipe4 requires the use of external programs, which can all be installed locally without root privileges (except Blast2GO which requires a MySQL database). The distribution comes with a set of helper scripts to automatically download and install all of the required software. All of the required programs are available under open-source licenses and are free to use (except Blast2GO which is not open source, but is free to use).

Results and Discussion

The analysis process

The above mentioned 9 steps can be described as follows (See Fig. 1 for a more graphical overview):

Step 1 – Extraction of the “FASTA” and “FASTA.QUAL” files from the original “SFF” file. This step can be skipped if not dealing with 454 data.

Step 2 – “Cleaning” the sequences, by discarding low overall quality and short reads, as well as reads that contain contaminants matched against the “UNIVEC” database [6] or any other contaminant database at the user’s discretion. This step uses the “Sequence Cleaner” program [7] and can also be skipped if dealing with Illumina data.

Step 3 – Assembling. This step uses mira [8]. A set of optimized parameters for SNP calling is contained in the example configuration file.

Steps 4 and 5 – SNP gathering. Resorting to the “MAF” output from step 3 (which is converted into the “SAM” [9] format), potential SNPs are identified in the assembly. The result is a summary intermediate “TCS” file and a “FASTA” file including all the “contigs” that contain putative SNPs (which are identified in the sequence title). The software “pysam” [9, 10] is used in this step.

Step 6 – Characterization of the detected SNPs, by attempting to fit them into Open Reading Frames (ORFs). The result is a “FASTA” file containing the ORFs with the SNPs identified in the sequence title, as well as the ORF frame allowing the quick assessment of the length and level of conservation of the SNP’s flanking region. This step uses the “EMBOSS getorf” program [11]. Also in this step, BLASTx [12] is run with the resulting ORFs against a large protein database, such as NCBI’s “nr”. Lastly, this step will produce an HTML formatted report with the
characterized SNPs for easy referencing. The report is formatted as a table and can easily be transferred to any spreadsheet software for further data exploring. Another output of this step is an additional HTML report with a compilation of various dataset metrics.
Step 7 (optional) – Blast2GO annotation; this step queries the contigs that contain SNPs against a large protein database such as NCBI’s 'nr' using BLASTx; these are then run through Blast2GO [13] using Blast2Go4Pipe, resulting in an annotation file that can be further analysed with Blast2GO itself.

Step 8 (optional) – SSR detection, by using “EMBOSS etandem” to detect potential SSRs in the assembly. The required quality of the putative SSRs is defined in the configuration file.

Step 9 – Compression of all the relevant result files into a 7zip archive which simplifies the transfer of (often large) results.

Example usage
A test dataset with documentation on example usage is provided with the software package. An example resulting report is also provided for the test dataset (run with default values on all settings).

Validation
In order to assess the efficiency of SNP detection and the rate of false positives, and assess the best default values to use, an approach using reference data was used.

For this goal, two reference sequences of two E. coli strains were used (http://www.ncbi.nlm.nih.gov/Traces/wgs/?val=ADWQ01 - Strain MS 85 and http://www.ncbi.nlm.nih.gov/Traces/wgs/?val=ADWR01 - Strain MS 79). Two 454 datasets were also downloaded from the NCBI Sequence Read Archive (SRA) [14] (http://www.ncbi.nlm.nih.gov/sra/SRX036805 and http://www.ncbi.nlm.nih.gov/sra/SRX036804) for the same strains as the references.

To assess the number of SNPs between the strains that could be found on the 454 datasets, the 454 reads of one strain were mapped against the reference sequences of the other strain using bowtie2 [15]. Atlas-SNP2 [16] reported 29673 SNPs between the reference sequence 85 and the 454 reads of the strain 79, and 28525 SNPs between the reference sequence 79 and the 454 reads of the strain 85.

4Pipe4 was then run on the two merged 454 datasets, discarding all strain information.

Although this validation method is not as good as true wet-lab genotyping, it is likely to be a good proxy, since Atlas-SNP2 is known to have very high sensitivity and specificity when dealing with 454 datasets [16].

The results varied with the different tested parameters (Table 1), but the best output was obtained with the default values of minimum coverage of 15 and minimum average quality of 70 per variant. This setup retrieved 114 SNPs, of which 32 did not match to any of those detected by Atlas-SNP2, being thus, considered false positives (28.07 % false discovery rate).

Although the number of provided SNPs is relatively low, due to the restrictive assembly and filtering parameters, we find this a good trade-off relative to the high confidence of the retrieved SNPs.

The task of SNP calling in 454 data has been performed before on organisms without a reference sequence or strain information, with varying degrees of false positives. One such study, conducted using custom scripts for SNP calling provided a false positive rate of 80 % on 4200 retrieved SNPs [17]. Another example, where the contigs of 283 SNPs were manually screened and selected, had a slightly better false positive rate of 45 % [18].

The above mentioned studies are not directly comparable to the results of the benchmark performed here, since they are performed on different datasets, nevertheless they can be used to infer that, in general, 4Pipe4 retrieves a smaller number of SNPs than other methods, but with a considerably lower false positive rate. Since the main goal of this pipeline is to provide the user with high confidence SNPs for genotyping arrays, a rate of 28.07 % false positives is a considerable improvement relative to the other mentioned approaches.

4Pipe4 compared to other software
Although 4Pipe4 is specifically designed for the purpose of detecting variation when no strain information or reference sequence is available, other software exists that can be used for the same purpose, but which differs from 4Pipe4 in some aspects:

QualitySNP [19] – Relies on CAP3 for clustering the reads (which is optimized for Sanger sequences, while 4Pipe4 uses mira, which is optimized for NGS data). Requires perl, PHP, a configured webserver and a MySQL

| Parameters used (Min. Average Quality) | 10/60 | 10/70 | 15/60 | 15/70 (Default) | 10/75 | 20/70 |
|---------------------------------------|-------|-------|-------|----------------|-------|-------|
| Total SNPs retrieved                  | 234   | 169   | 155   | 114            | 107   | 89    |
| Confirmed SNPs                        | 97    | 86    | 88    | 82             | 69    | 57    |
| False Positive rate (%)               | 58.55 | 49.11 | 43.23 | 28.07          | 35.51 | 35.96 |
database for SNP retrieval. This means that root access to the machine in which the program is being run on is required. Furthermore, QualitySNP has been superseded by the simpler and faster QualitySNPng [20].

AGSNP [21] – Relies on Newbler assembler for clustering, and if strain information is not available, it further requires combining 454 data with Illumina or SOliD data (4Pipe4 does not require multiple technologies data for SNP calling).

Still other programs exist for SNP detection, but they usually require either a reference sequence, such as Atlas-SNP2 or SAMtools, or strain information, such as discoSnp++ [22] (formerly kisSnp [3]) or DIAL [23].

There is, however, another program that can be used for the same purpose as 4Pipe4 – QualitySNPng. This program, however, is not an analysis pipeline, but rather a SNP caller for read alignments. It has a graphical user interface, which can be disabled for use in servers, but still requires “Qt4” to be installed in the server (which is not frequent). In order to compare it with 4Pipe4, we have modified the program to be usable without “Qt4” installed (https://github.com/StuntsPT/4Pipe4) and provide a branch of 4Pipe4 which is ready to use QualitySNPng (https://github.com/StuntsPT/4Pipe4/tree/new_snп_caller), without requiring any further dependencies.

Benchmarking the results of 4Pipe4 with QualitySNPng as the SNP caller, more SNPs were returned (513/147 SNPs found with the default/tuned values) than with our SNP caller, but with a larger rate of false positives (only 60,22 SNPs were a match to those found by AtlasSNP2, meaning a false positive rate of 88.3 %|85 %). Therefore, the built-in SNP caller was kept as default, but QualitySNPng can still be used from its own git branch if desired.

For the sake of completeness, we also made the SNP calling on the benchmark dataset using the software discoSnp++ (which requires strain identification) with the most restrictive parameters, to minimize the number of false positives. This program retrieved 9226 SNPs, of which 5967 were considered true positives (false positive rate of 35.3 %). As expected, this method retrieves more SNPs than both 4Pipe4 and QualitySNPng since it takes advantage of strain information, but it still provides a somewhat higher false positive rate than 4Pipe4.

Conclusions

We present here an automated analysis process specifically designed for SNP detection from 454 pyrosequencing transcriptome reads, which we named 4Pipe4. This is the first program specifically built to automate the whole process of finding putative SNPs in NGS datasets that lack both information regarding the origin of each read and a reference sequence. In-silico validation of 4Pipe4 results using previously analysed reference data revealed good performance in the calling of high confidence SNPs.

The 4Pipe4 pipeline, at the cost of retrieving a relatively low number of SNPs, has provided a lower rate of false positive SNPs than both an alternative SNP caller (QualitySNPng) and an alternative software that uses strain information (discoSnp++), as well as those obtained in previous studies that used different approaches for a similar type of data and goal.

Since the main purpose of this software is to retrieve high confidence SNPs for further exploring, we expect the incremental contributions it brings to improve, speed up and facilitate research on the field of population genomics.

Furthermore, we expect to implement new features in 4Pipe4, such as: graphics in the metrics report; indel variation finding; integration of alternative software (such as newbler for assembling instead of mira); process optimization for NGS technologies besides 454; switch from FASTA + FASTA.QUAL format to FASTQ. These are some of the planned features, but others can be requested and implemented, should there be demand for them.

Availability and requirements

- Project name: 4Pipe4
- Project home page: https://github.com/StuntsPT/4Pipe4
- Operating system(s): Platform independent
- Programming language: Python
- Other requirements: Python 3.0 or higher for running the pipeline, several other programs for specific tasks (consult the README file)
- License: GNU GPLv3
- Any restrictions to use by non-academics: None

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

FPM has developed the software, drafted the SNP detection routines and written the manuscript; BMV has extensively reviewed the code and assisted in the initial data analyses; SGS has provided valuable insights on SNP data interpretation and proofread the manuscript; DB has provided the samples and the datasets for the initial data analyses, and proofread the manuscript and OSP has conceived of the study, and participated in its design and coordination.

Acknowledgements

This work was fully supported by projects SOBEREIRO/0036/2009 (under the framework of the Cork Oak ESTs Consortium), PTDC/BIA-BEC/098783/2008 and PTDC/AGR-GPL/119943/2010 from Fundação para a Ciência e Tecnologia (FCT) – Portugal. F. Pina-Martins was funded by FCT grant SFRH/BD/51411/2011, under the PhD program “Biology and Ecology of Global Changes”, Univ. Aveiro & Univ. Lisbon, Portugal. D. Batista was funded by FCT grant SFRH/BPD/104629/2014. We would furthermore like to thank the...
reviewers of the manuscript, whom have provided very constructive criticism, resulting in great improvements to both the manuscript and the software.

**Author details**
1. Departamento de Biologia Animal, Faculdade de Ciências, Computational Biology and Population Genomics Group, cEB - Centre for Ecology, Evolution and Environmental Changes, Universidade de Lisboa, Campo Grande, 1749-016 Lisboa, Portugal. 2. Departamento de Biologia e CESAM, Univ. de Aveiro, Aveiro, Portugal. 3. Wurm Lab, School of Biological & Chemical Sciences, Queen Mary University of London, Mile End Road, E1 4NS London, UK. 4. Centro de Investigação das Ferrugens do Cafeeiro (CIFC), Instituto Superior de Agronomia (ISA), Universidade de Lisboa, Quinta do Marquês, 2784-505 Olris, Portugal.

**Received:** 31 March 2015 **Accepted:** 15 January 2016

---

**References**

1. Schuster SC. Next-generation sequencing transforms today's biology. Nat Methods. 2008;5:16–8.

2. Papanicolaou A, Sterli R, French-Constant RH, Heckel DG. Next generation transcriptomes for next generation genomes using est2assembly. BMC Bioinformatics. 2009;10:447.

3. Peterlongo P, Schnel N, Pisanti N, Sagot MF, Lacroix V. Identifying SNPs without a Reference Genome by Comparing Raw Reads. In: Chaves E, Lioradi S, editors. String Processing and Information Retrieval. Springer Berlin Heidelberg, 2010. p. 147–58.

4. Modesto IS, Miguel C, Pina-Martins F, Glushkova M, Veloso M, Paula OS, et al. Identifying signatures of natural selection in cork oak (Quercus suber L.) genes through SNP analysis. Tree Genet Genomes. 2014;10:1645–60.

5. Savage AE, Kiemer-Tabuczy KM, Ellison AR, Fleischer RC, Zamudio KR. Conservation and divergence in the frog immunome: pyrosequencing and de novo assembly of immune tissue transcriptomes. Gene. 2014;542:98–108.

6. The UniVec Database [http://www.ncbi.nlm.nih.gov/tools/vecscreen/univec/]. Accessed 12 Jan 2012.

7. Sequence Cleaner [http://sourceforge.net/projects/seqclean/]. Accessed 10 Jan 2012.

8. Chevreux B, Pfisterer T, Drescher B, Driesel A, Muller WEG, Wetter T, et al. Using the miraEST assembler for reliable and automated mRNA transcript assembly and SNP detection in sequenced ESTs. Genome Res. 2004;14:147–59.

9. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, et al. The Sequence Alignment/Map format and SAMtools. Bioinforma Oxf Engl. 2009;25:2078–9.

10. pysam-developers/pysam [https://github.com/pysam-developers/pysam]. Accessed 01 Mar 2014.

11. Rice P, Longden I, Bleasby A. EMBOSS: The European molecular biology open software suite. Trends Genet TIG. 2000;16:276–7.

12. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990;215:403–8.

13. Conesa A, Gotz S, Garcia-Gomez JM, Terol J, Talon M, Robles M. Blast2GO: a universal tool for annotation, visualization and analysis in functional genomics research. Bioinforma Oxf Engl. 2005;21:3674–6.

14. Leinonen R, Sugawara H, Shumway M. The sequence read archive. Nucleic Acids Res. 2011;39(Database issue):D19–21.

15. Langmead B, Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nat Methods. 2012;9:357–9.

16. Shen Y, Wan Z, Coara C, Dabrek R, Chen L, Ostrowski EA, et al. A SNP discovery method to assess variant allele probability from next-generation resequencing data. Genome Res. 2010;20:273–80.

17. Tollaenare C, Susi H, Noko-Kovisto J, Koskinnen P, Tack A, Auvinen P, et al. SNP Design from 454 Sequencing of Podosphaera plantaginis Transcriptome Reveals a Genetically Diverse Pathogen Metapopulation with High Levels of Mixed-Genotype Infection. PLoS One. 2012;7:e52492.

18. Broders KD, Woeste KE, San Miguel PJ, Westernman RF, Boland GJ. Discovery of single-nucleotide polymorphisms (SNPs) in the uncharacterized genome of the ascomycete Ophiognomonia clavigignenti-juglandacearum from 454 sequence data. Mol Ecol Resour. 2011;11:693–702.

19. Tang J, Vosman B, Voorrips RE, van der Linden CG, Unessnism JAM. QualitySNP: a pipeline for detecting single nucleotide polymorphisms and insertions/deletions in EST data from diploid and polyploid species. BMC Bioinformatics. 2006;7:438.

20. Njooev H, van Kaauwen M, Esselink DG, Hoegen B, Vosman B. QualitySNPng: a user-friendly SNP detection and visualization tool. Nucleic Acids Res. 2013;41:W587–90.

21. You FM, Huo N, Deal KR, Gu YQ, Luo M-C, McGuire PE, et al. Annotation-based genome-wide SNP discovery in the large and complex Aegilops tauschii genome using next-generation sequencing without a reference genome sequence. BMC Genomics. 2011;12:59.

22. Uricaru R, Rok G, Lacroix V, Quillery E, Plantard O, Chikhi R, et al. Reference-free detection of isolated SNPs. Nucleic Acids Res. 2014;43:e11.

23. Ratan A, Zhang Y, Hayes VM, Schuster SC, Miller W. Calling SNPs without a reference sequence. BMC Bioinformatics. 2010;11:90.

---

Submit your next manuscript to BioMed Central and we will help you at every step:

- We accept pre-submission inquiries
- Our selector tool helps you to find the most relevant journal
- We provide round the clock customer support
- Convenient online submission
- Thorough peer review
- Inclusion in PubMed and all major indexing services
- Maximum visibility for your research

Submit your manuscript at www.biomedcentral.com/submit