

User Manual

User manual version 3.1

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For the latest version of SPANDx and its associated user manual, please visit either of the following SPANDx websites: http://sourceforge.net/projects/spandx/ or https://github.com/dsarov/SPANDx

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INTRODUCTION AND DESCRIPTION

SPANDx (<u>S</u>ynergised <u>P</u>ipeline for <u>A</u>nalysis of <u>N</u>ext-generation sequencing <u>D</u>ata in Linu<u>x</u>) is a comparative genomics pipeline designed to greatly simplify the identification of genetic variants (i.e. single-nucleotide polymorphisms [SNPs], insertions/deletions [indels], and large [>100bp] deletions) from medium- to large-sized haploid next-generation re-sequencing (NGS) datasets. SPANDx can process several NGS data inputs including paired- and single-end data from the Illumina MiSeq, HiSeq and GA_{IIx} platforms, single-end data from the Life Technologies Ion Personal Genome Machine (PGM)[®], and single-end Roche 454 data. SPANDx integrates the following validated bioinformatics tools for start-to-finish sequence analysis of raw NGS data using a single command:

- Burrows-Wheeler Aligner (BWA) (1, 2) for alignment of short (i.e. Illumina and PGM) or long (i.e. 454) NGS read data. BWA is downloadable from http://bio-bwa.sourceforge.net/.
 SPANDx does not currently support BWA versions later than 0.6.2.
- **SAMtools** (3) and **Picard** (as-yet-unpublished) for alignment manipulation and filtering. These programs can be downloaded from http://samtools.sourceforge.net/ and http://broadinstitute.github.io/picard/, respectively.
- **Genome Analysis Tool Kit (GATK)** (4-6) for base quality score recalibration, realignment of regions with low mapping quality, duplicate removal, identification of SNPs and indels, and filtering the variant call format (VCF) file generated from the alignment process. GATK can be downloaded from http://www.broadinstitute.org/gatk/.
- VCFtools (7) for manipulation of VCF files. VCFtools is downloadable from https://vcftools.github.io/index.html. All file outputs from SPANDx are in standardised VCFv4.1 format. tabix and bgzip are VCFtools dependences required for handling.vcf files, and can be downloaded here: http://sourceforge.net/projects/samtools/files/tabix/.
- **BEDTools** (8), and specifically the *coverageBED* module, for identification of locus presence/absence across each genome of interest based on the reference sequence. This tool is useful for identifying larger-scale (approx. 100bp or larger) deletions. BEDTools can be downloaded from https://github.com/arq5x/bedtools2/.
- **SnpEff** (9) for annotation of SNP and indel variants. SnpEff can be downloaded from here: http://snpeff.sourceforge.net/.
- **PLINK** (10) for microbial genome-wide association studies (mGWAS). PLINK can be downloaded from http://pngu.mgh.harvard.edu/~purcell/plink/download.shtml.

All of the above dependencies, with the exception of GATK and PLINK, come pre-bundled and pre-compiled with the SPANDx package. GATK needs to be downloaded and installed separately due to Broad Institute licencing restrictions. PLINK has not been included in the SPANDx bundle yet, but will be included in future versions.

The dependency binaries have been compiled for an x86-64 Linux system. If you have different system architecture, you will need to install SPANDx dependencies yourself.

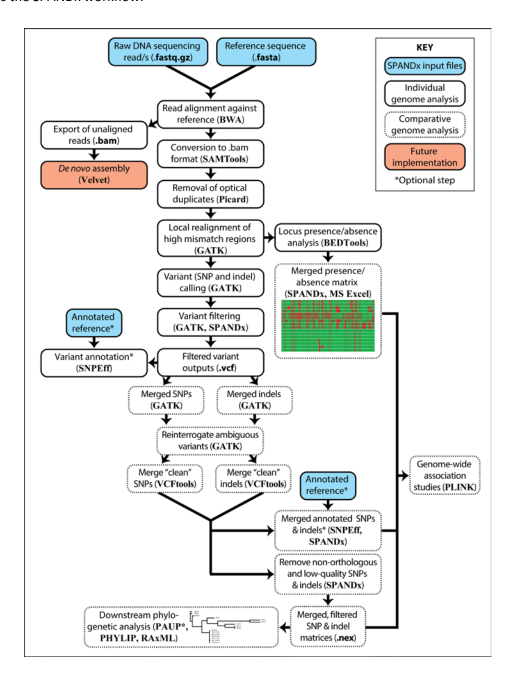
Novel comparative genomic features of SPANDx include:

- <u>Merged orthologous¹ SNP and indel matrices</u> that greatly simplify variant visualisation for comparative genomic analyses.
- PAUP*, PHYLIP and RAxML-compatible .nex core genome orthologous¹ SNP and indel matrix files for downstream phylogenetic analyses. PHYLIP and RAxML are freely available programs downloadable from: http://evolution.genetics.washington.edu/phylip.html and https://github.com/stamatak/standard-RAxML. As of July 2015, PAUP* is now open source and apparently will be updated by the developer over the coming months. Please use Google to find the latest version!
- <u>Locus presence/absence matrices</u> from BEDTools outputs that enable simple visualisation and comparative genomic determination of 1) the core genome and 2) variable genetic loci (including deleted regions brought about by e.g. reductive evolution).
- <u>Merged, annotated SNP and indel matrices</u> for fast and simple genetic characterisation of variants (NB. The user must provide SPANDx with the SnpEff-annotated reference genome information for variant annotation to work).

¹SNP and indel matrices generated by SPANDx exclude variants in paralogous loci, as well as variants that reside in genetic loci missing in one or more genomes. If these variants are required, they can be found in the individual filtered .vcf output files. Alternatively, all variants are outputted in the annotated matrix file/s.

Unlike many other comparative bioinformatic tools, SPANDx does not require pre-assembled genomes. The major advantage of this approach is that the raw NGS reads are used for variant detection, resulting in a low probability of error and a high variant detection rate. In addition, the default settings for variant calling using Illumina, Ion PGM and 454 data have already been optimised in SPANDx and do not require the user to specify these settings, although these settings can be customised if required.

SPANDx has been purposely written for systems that utilise Portable Batch System (PBS), Sun Grid Engine (SGE) [i.e. qsub] or Simple Linux Utility for Resource Management (SLURM) [i.e. sbatch] to enable task parallelisation, greatly reducing turn-around-time of datasets comprising tens through to thousands of haploid genomes using a single command. As of version 2.7, SPANDx will also run on systems without a resource manager but will not run jobs in parallel. Currently there is no support for SPANDx on other resource management systems (e.g. LSF) due to the unavailability of such systems in our laboratory, but compatibility with resource managers can be addressed by the developers if required. Please email us at mshr.bioinformatics@gmail.com if you require a specific resource manager compatible version of SPANDx and are willing to test it on your system.



SYNOPSIS

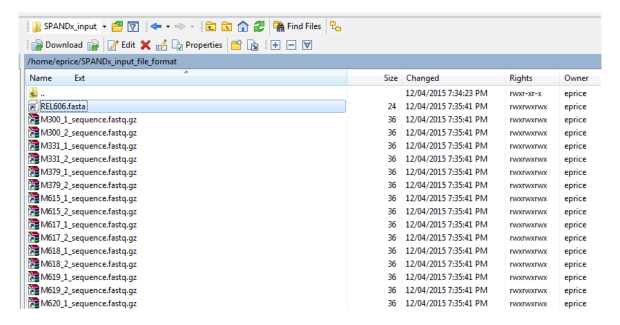
SPANDx -r <exact reference name, excluding .fasta extension> [parameters, optional] -o [organism] -m [generate SNP matrix yes/no] -i [generate indel matrix yes/no] -a [include annotation yes/no] -v [reference file for variant annotation; name must exactly match the SnpEff database name, which can be found in the snpEff.config file] -s [specify read prefix to run single strain; set to none to construct a SNP matrix from a previous analysis or leave as default to process all reads] -t [sequencing platform i.e. Illumina/Illumina_old/454/PGM] -p [pairing of reads (i.e. paired-end or single-end) PE/SE] -w [BEDTools window size in base pairs] -z [include tri-allelic and tetra-allelic SNPs yes/no]

COMMANDS AND OPTIONS

SPANDx.sh is the only script that needs to be run to obtain data outputs. SPANDx by default expects paired-end Illumina data with v1.8+ quality encoding. If your data are in this format, the only required switch is $\neg x$ to specify the reference sequence prefix. If another NGS data format is being analysed, please specify this format using the $\neg t$ (and if single-end, the $\neg p$) switch/es². By default, SPANDx <u>will</u> construct a locus presence/absence matrix but <u>will not</u> construct orthologous SNP or indel matrices, nor will it perform variant annotation. The $\neg m$ (<u>matrix</u>) and $\neg t$ (<u>indel</u>) switches are required for core genome orthologous SNP and indel matrix creation, respectively. The $\neg a$ (<u>a</u>nnotate) and $\neg v$ (reference name for <u>v</u>ariant identification) switches are both required for variant annotation.

²SPANDx cannot process multiple NGS formats (e.g. single-end and paired-end Illumina) in a single run. If multiple NGS formats are to be analysed, please create separate analysis directories and run SPANDx specifically for each NGS format. These data can be merged for downstream analysis. See the -s description for more information.

Prior to running SPANDx.sh, both the reference (in .fasta format) and NGS files (in .fastq.gz format) are required to be in your analysis directory. SPANDx expects all NGS reads to be in the sequence analysis directory (i.e. the present working directory) and by default all NGS reads within the sequence analysis directory will be processed. Before running SPANDx, make sure the NGS read files conform to the following format regardless of the sequencing technology used: strain_1_sequence.fastq.gz and strain_2_sequence.fastq.gz (for paired-end reads) or strain_1_sequence.fastq.gz (for single-end reads). See screenshot below for correctly formatted reference genome and paired-end input files.



Options:

- -r STR Required. Specifies the reference genome file, excluding the .fasta extension. The -r switch is the only mandatory switch needed for SPANDx to function. Additional switches are required to modify the default behaviour of SPANDx. Sequencing technology needs to be specified if your data are not paired-end Illumina data with v1.8+ quality scores. The reference file is required to be in .fasta format and should conform to the standard FASTA specification, as detailed here: http://www.ncbi.nlm.nih.gov/BLAST/blastcgihelp.shtml. IUPAC codes are not supported by some programs incorporated in SPANDx and must be avoided. In addition, the .fasta reference file must not contain any blank lines. For compatibility with the annotation module of SPANDx, the chromosome names for the reference genome must match those used by SnpEff. This nomenclature can be found in the snpEff.config file, which is generated upon SnpEff installation, or automatically with the full SPANDx installation.
- -o *STR Optional.* Specifies the organism under analysis. The -o parameter is used in naming the read group headers in the SAM and BAM files. Spaces and special characters may have unexpected behaviour and should be avoided. [Haploid]
- -m yes/no Optional. The -m switch is used to create a matrix with all orthologous SNP variants identified in the analysis. Non-orthologous SNPs are excluded. Output .nex files can be directly imported into PAUP*, PHYLIP or RAxML for downstream phylogenetic analysis. By default this behaviour is switched off. [no]
- -i yes/no Optional. The -i switch is used to create a matrix with all orthologous indel variants identified in the analysis. Non-orthologous indels are excluded. By default this behaviour is switched off. [no]
- -a yes/no Optional. The -a switch is used to perform annotation of the variant files. By default this behaviour is switched off. If annotation is switched on the -v switch must also be specified. [no]
- -v STR Optional, required if -a is set to "yes". The -v switch is used to specify the reference genome that SnpEff will use to annotate variants. The string used for this variable must match one of the genomes contained within the SnpEff.config file. Additionally, chromosome names in the reference file must match those contained within the SnpEff program. Please refer to the SnpEff manual (which can be found here: http://snpeff.sourceforge.net/SnpEff manual.html) if you require more information. [null]
- optional. The -s switch is used to flag a single genome for analysis. If -s is set to none, SPANDx will not perform individual analysis of any NGS read files in the current directory. Instead, SPANDx will move to the comparative genomics section of the pipeline (see SPANDx workflow above) and assume all individual genome analysis has already been completed. SPANDx will then merge all VCF files contained in \$PWD/Phylo/snps and perform error correction using the .bam and .bai files contained in \$PWD/Phylo/bams to construct an orthologous SNP matrix, which will be output to \$PWD/Phylo/out. Before running this module, please check that all .vcf files located within \$PWD/Phylo/snps match the alignment files within \$PWD/Phylo/bams, and

that all .bam files contain their accompanying .bai index file. This feature mitigates the need to rerun previous analyses from scratch, and is useful for combining data generated from multiple SPANDx runs (e.g. from different sequencing technologies) into a single orthologous SNP matrix. [all]

- -t STR Optional. The -t switch specifies the sequence technology used, and must be one of the following: Illumina, Illumina_old, 454 or PGM. By default, SPANDx expects Illumina reads with Phred+33 read quality encoding, which is standard as of v1.8+. To specify NGS reads generated by the older Illumina format (i.e. Phred+64), use -t Illumina_old. If the analysis mode is switched to 454 or PGM, SPANDx will use the BWA-SW algorithm of BWA for read alignment and thus will expect reads to be in single-end strain_1_sequence.fastq.gz format with Phred+33 quality encoding. [Illumina]
- -p STR Optional. The -p switch specifies the pairing of reads and must be either PE or SE. By default SPANDx expects reads to be paired. If reads are single end, -p must be set to SE. Currently SPANDx does not support paired-end 454 or PGM data. [PE]
- $_{-W}$ INT Optional. The $_{-W}$ switch specifies the window size (in base pairs) used by BEDTools to analyse whole genome alignment coverage i.e. locus presence/absence. [1000]
- -z yes/no Optional. The -z switch enables users to output tri- and tetra-allelic SNPs in their SNP matrix output. By default, SPANDx will only output biallelic SNPs. [no]

INSTALLATION AND REQUIREMENTS

SPANDx is written in Bash and will run on most Linux installations. For parallelisation, SPANDx can utilise PBS, SGE or SLURM resource managers. As of version 2.7, SPANDx will also run on systems without a resource manager, but will not run in parallel. SPANDx has been tested on GNU Bash version 3.2.25(1)-release and GNU Bash version 4.1.2(1)-release (both on x86_64-redhat-linux-gnu) with Java v1.7.0_55 and v1.7.0_71. SPANDx v3.0 (which is downloadable here: http://sourceforge.net/projects/spandx/ or here: https://github.com/dsarov/SPANDx) has been tested using PBS (TORQUE v2.5.13), SGE v6.2u5p3, and SLURM v14.11. Tested dependency versions are BWA v0.6.2-r126, SAMtools v0.1.18, v0.1.19 and 1.2, Picard v1.134, the Genome Analysis Toolkit v3.2.2, BEDTools v2.18.2, SnpEff v4.1 (build 2015-01-07), VCFtools v0.1.11 and tabix v0.2.5 (r1005). SPANDx does not currently support BWA versions later than 0.6.2.

For parallelisation, <u>SPANDx</u> requires <u>PBS</u>, <u>SGE</u> or <u>SLURM</u> to submit jobs to the cluster. If you do not have one of these system setups, SPANDx cannot be run in parallel. Please contact us if you are wanting to trial SPANDx on your system but are using a different Linux resource scheduler setup than those listed here.

SPANDx should work from any installation directory but has been most extensively tested in /home/user/bin. To install SPANDx, gunzip and untar the program (usually with the command tar xvfz SPANDx_vx.x.tar.gz) in your /bin directory. Optionally, the latest version of SPANDx can be downloaded from GitHub page using the following command: git clone https://github.com/dsarov/SPANDx. If you download from GitHub, you may need to change file permissions prior to running.

IMPORTANT: following extraction of the script files, the SPANDx.config file will need to be modified to contain the location of the SPANDx installation. If your system uses a proxy to access the internet, please modify the JAVA_PROXY variable in SPANDx.config. In addition, you will also need to download and install GATK and either place the GenomeAnalysisTK.jar file (renamed without version numbers) in the SPANDx installation directory or specify the install path in the SPANDx.config file.

For users who <u>don't</u> have x86-64 systems: You will need to download and install SPANDx dependencies yourself as only x86-64 binaries are included in the SPANDx distribution. Following extraction of the script files, the <u>SPANDx.config</u> file will need to be modified to contain the absolute paths of each dependency (i.e. BWA, SAMTools, Picard, GATK, BEDTools, SnpEff, VCFtools and tabix installations – see Introduction and Description above for web links to these free third-party programs). If a "dependency not found" error occurs, please check the installation path and the location specified in the <u>SPANDx.config</u> file.

Please make sure to specify the location of the PERL5 libraries (automatically installed with VCFtools) in the SPANDx.config file. PERL5 libraries are required for correct functioning of VCFTools. Make sure that the location of tabix and bgzip (dependencies of VCFtools) are specified in your PATH variable.

Some of these programs have additional dependencies that are required for them to function properly e.g. Java. Please refer to the appropriate manuals or system administrator for installation of these utilities if they are not already on your system.

SPANDx customisation:

Resource manager:

Depending on your cluster environment, the scheduler.config file may need to be changed. By default, SPANDx will expect the resource manager to be PBS. IMPORTANT: If you are using SGE, please modify the SCHEDULER variable to SGE. If you are using SLURM, please modify the SCHEDULER variable to SLURM. If no resource manager is available, please modify the SCHEDULER variable to NONE. This file will configure the operation of the resource manager. By default, all commands request one node and 12 hours of wall time, you will not be sent mail, and standard output is merged with standard error. These settings can be changed if your job needs more time to complete or if you want e-mail notifications of job completion.

Variant calling:

One advantage of SPANDx over other tools is that GATK variant calling parameters are already specified. These parameters have been tested across NGS data for several bacterial species generated on different NGS platforms. Therefore, the default settings should work well for most haploid genome projects. If desired, users can customise the SPANDx variant filtering parameters by altering the GATK.config file. All filtering steps used in GATK can be customised using this configuration file. Note that these variables must conform to JEXL specifications.

SPANDx variant calling has been optimised for bacterial genomes so may behave differently for other haploid organisms. If in doubt, outputs should be verified with e.g. wet lab analysis of variants. SPANDx is currently not configured to analyse diploid or polyploid genomes.

The following parameters can be customised to change the variant filtering behaviour for both SNPs and indels if required; below are the default SPANDx settings:

```
CLUSTER_SNP=3 (for SNPs only)
CLUSTER_WINDOW_SNP=10 (for SNPs only)
MLEAF=0.95
QD=10.0
MQ=30.0
FS=10.0
HAPLO=20.0
QUAL=30.0
```

LOW_DEPTH=2 (variants with less than the average coverage, divided by LOW_DEPTH, will fail filtering. If this value is set at the default of 2, regions with less than half the average depth will fail and thus will be filtered out).

HIGH_DEPTH=3 (a value of 3 means that regions with more than three times the average coverage of the entire genome will fail and thus will be filtered out).

INTERPRETING THE OUTPUTS

Summary statistics:

As of v3.1, SPANDx provides a summary of all SNPs, indels and X coverage in \$PWD/Outputs/Single_sample_summary.txt.

Alignment files:

Alignment files generated with SPANDx are in .bam format and can be found in \$PWD/strain/unique/strain.realigned.bam. If visualisation of the alignment is desired, these files can be easily viewed in an alignment viewer (our favourite is Tablet (11); downloadable from: http://ics.hutton.ac.uk/tablet/).

SNP and indel variants:

SNPs and indels are output into two locations: 1) <code>\$PWD/Outputs/SNPs_indels_PASS</code>, which contains both SNP and indel variants that have passed the filters specified in the <code>GATK.config</code> file (see "Variant calling" above for details of the default filters); and 2) <code>\$PWD/Outputs/SNPs_indels_FAIL</code>, which contains SNPs and indels that have failed filtering parameters.

SPANDx can repeat variant filtering steps without also repeating the alignment and data processing steps. To use this behaviour, remove the relevant <code>snps.PASS</code> and <code>indels.PASS</code> files from the <code>\$PWD/Outputs/SNPs_indels_PASS</code> directory and the relevant <code>snps.AMB</code> and <code>indel.AMB</code> files from the <code>\$PWD/Outputs/SNPs_indels_FAIL</code> directory, change the <code>GATK.config</code> file to the desired parameters, and re-run SPANDx. NB. SPANDx will only re-filter the variants with altered parameters for those strain/s that have been removed from the <code>Output</code> directories.

Whole genome coverage (a.k.a. locus presence/absence):

Following assessment of whole genome coverage by BEDTools (8), SPANDx provides a combined BEDcov matrix for all analysed genomes in: \$PWD/Outputs/Comparative/Bedcov_merge.txt. This file lists the BEDTools 'windows', or NGS read coverage, for each strain based on the reference sequence, and ranges from 0 (0% read coverage across the window) to 1 (100% coverage across the window). The BEDcov matrix can be imported into Microsoft Excel for easier visualisation and manipulation. Bedcov_merge.txt is a useful file for identifying the core genome of a given dataset and for identifying variable genetic regions among strains. The base-pair resolution of this output can be modified by changing the -w switch (the default in SPANDx is 1000). We recommend changing to 100 for microbial GWAS analysis in PLINK to increase sensitivity, and leaving as default for all other analyses.

Orthologous SNP matrices for phylogenetic analyses:

Two separate SNP matrix files are generated by SPANDx. These matrices are output in \$PWD/Outputs/Comparative/ and are named Ortho_SNP_matrix_RAxML.nex and Ortho_SNP_matrix.nex.

The Ortho_SNP_matrix files generated by SPANDx exclude SNPs that are low-quality, that are in non-orthologous regions, and that are tri- or tetra-allelic in nature. Non-orthologous SNPs should not be used for phylogenetic reconstruction. In addition, filtering for tri- and tetra-allelic SNPs is performed to minimise erroneous calls (which can look like tri- and tetra-allelic SNPs) passing through filters.

NB. The annotated SNP matrix (All_SNPs_annotated.txt) include tri- and tetra-allelic variants. In the annotated SNP and indel matrices, ambiguous and non-orthologous calls are flagged as "?" or ".", respectively.

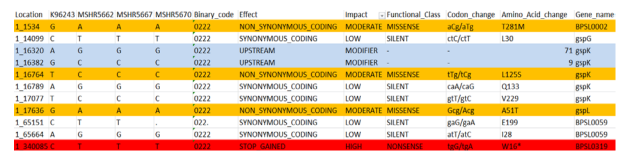
Ortho_SNP_matrix.nex includes SNP coordinates identified by GATK. This file is directly importable into PAUP* and is useful for phylogenetic estimations that require nucleotide data (e.g. maximum likelihood). Below is a screen capture of this file:

Ortho_SNP_matrix_RAxML.nex is a RAxML- and PHYLIP-importable version of the Ortho_SNP_matrix_RAxML.nex file. Note that for compatibility with PHYLIP, taxa names must have exactly 10 characters (including spaces) otherwise Ortho_SNP_matrix_RAxML.nex will not be recognised as a valid PHYLIP file. SPANDx does not automatically rename taxa to meet this PHYLIP requirement.

Annotation:

When annotation is switched on, annotated SNP and indel variants will be generated for each genome under analysis; these variants will be output to \$PWD/strain/unique/annotated. If both annotation and comparative analysis is switched on (-a yes and -m yes), annotated, merged SNP and indel matrices will be generated for all genomes under analysis. These files can be found in \$PWD/Outputs/Comparative and are called All_indels_annotated.txt and All_SNPs_annotated.txt. These tab-delimited text files can be easily imported into Excel, as

shown in the screenshot below. Note that the Binary_code column will need to be specified as "Text only" due to the character string containing "0". This column is a representation of the SNP/indel pattern across all genomes for each variant, and can be useful for filtering purposes.



What do the codes in the annotated matrices mean?

<u>SNPs</u>: **0** matches the reference genome. **2**, **3** and **4** represent bi-, tri- and tetra-allelic variants. **?** represents low quality (ambiguous) SNP calls; these variants should be interpreted with caution.

<u>Indels</u>: **0** represents the reference genome; as of SPANDx v3.0, **2** thru **9** represent variants. Indel loci with ≥10 variants are not currently identified in the annotated matrix, so variants exceeding this number will be tagged by **?**. As with SNPs, **?** can also represent low quality.

<u>SNPs & indels</u>: SNP/s indels represented by a full stop (.) do not pass the depth filters and are either in absent (e.g. deleted) or in very low-coverage regions.

On a somewhat tangential note, we have found that the All_SNPs_annotated.txt is extremely helpful for detecting mixtures in NGS reads. If you see one or more genomes with unusually high prevalence of ?, it is probably mixed!

Merged SNP-indel matrices:

In certain instances, particularly in outbreak studies where only closely related strains are being examined and there are few genetic variants, it may be desirable to merge SNP and indel variants prior to phylogenetic analysis for increased resolution. We have recently shown that this approach can provide more robust phylogenetic correlation with epidemiological data than constructing a phylogeny based on SNPs alone (12). To merge SNP and indel variants into a separate output file, run the MergeSnpIndel.sh script while in the \$PWD/Outputs/Comparative/ directory. The MergeSnpIndel.sh script will merge and reformat the data in the Ortho_SNP_matrix.nex and indel_matrix.nex files. The output file is: indel_SNP_matrix.nex. This file is directly importable into PAUP*. Below is a screen capture of this file:

Genome-wide association studies (GWAS):

The main comparative outputs of SPANDx (SNP matrix, indel matrix, presence/absence matrix, annotated SNP matrix and annotated indel matrix in \$PWD/Outputs/Comparative/) can be used as input files for GWAS. From version 2.6 onwards, SPANDx is distributed with GeneratePlink.sh.

The GeneratePlink.sh script requires two input files: an ingroup.txt file and an outgroup.txt file. The ingroup.txt file should contain a list of the taxa of interest (e.g. antibiotic-resistant strains) and the outgroup.txt file should contain a list of all taxa lacking the genotype or phenotype of interest (e.g. antibiotic-sensitive strains). The ingroup.txt and outgroup.txt files <u>must</u> include only one strain per line. Although larger taxon numbers in the ingroup and outgroup files will increase the statistical power of GWAS, it is better to only include relevant taxa i.e. do not include taxa that have not yet been characterised, or that have equivocal data. The GeneratePlink.sh script will generate .ped and .map files for SNPs, and presence/absence loci and indels if these were identified in the initial analyses. The .ped and .map files can be directly imported into PLINK. For more information on GWAS and how to run PLINK, please refer to the PLINK website: http://pngu.mgh.harvard.edu/~purcell/plink/

Log files:

By default, both the standard error and standard output are merged into a single log file. Almost all commands in SPANDx are captured in log files. If an error occurs, the log files are a good first place to look. As of SPANDx 3.0, those using a PBS scheduler can find their log files here: \$PWD/logs. For everyone else, log files can be found in \$PWD.

If you wish to minimise the amount of log files that SPANDx generates, you can change the ERROR_OUTPUT variable (PBS) to n, or the ERROR_OUTPUT_SGE (SGE) to no, in the scheduler.config file. Please note that this feature only works when using PBS or SGE resource handlers.

Unaligned reads:

A .bam file of the unaligned reads is generated by SAMTools after BWA alignment. The unaligned reads can be found in: \$PWD/strain/unique/unmapped.bam. It is anticipated that future versions of SPANDx will include an option for automated assembly of these unaligned reads.

EXAMPLES

The simplest way to run SPANDx is if your reads are in paired-end, Illumina format and follow the naming convention of strain_1_sequence.fastq.gz and strain_2_sequence.fastq.gz. SPANDx can then be run by simply specifying the reference .fasta genome prefix. All read files in the current directory will be processed, although a SNP or indel matrix will not be constructed nor will variant annotation be performed unless specified.

"No frills" SPANDx command for basic Illumina 1.8+ analysis:

```
/home/user/bin/SPANDx/SPANDx.sh -r reference (without .fasta extension)
```

If other SPANDx features are required or reads other than Illumina v1.8+ are used, these features will need to be specified as per the examples below.

Paired-end Illumina 1.8+ reads, SNP matrix required, no indels required, no annotated genome available/required:

```
/home/user/bin/SPANDx/SPANDx.sh -r REL606 -m yes
```

To include an annotation for the above example:

```
/home/user/bin/SPANDx/SPANDx.sh -r REL606 -a yes -m yes -v Escherichia coli B REL606 uid58803
```

Paired-end Illumina 1.8+ reads, indel matrix required, no annotated genome available/required:

```
/home/user/bin/SPANDx/SPANDx.sh -r REL606 -i yes -m yes
```

Paired-end Illumina 1.3 reads, SNP and indel matrices required, annotated reference genome Hi_86-028NP.fasta available:

```
/home/user/bin/SPANDx/SPANDx.sh -r Hi_86-028NP -o Hi -m yes -i yes -a yes -v Haemophilus_influenzae_86_028NP_uid58093 -t Illumina_old
```

Single-end Ion PGM reads, SNP/indel matrices and annotation not required, BEDCov window size of 500bp (instead of the default 1000bp) desired, using the same reference genome as above:

```
/home/user/bin/SPANDx/SPANDx.sh -r Hi_86-028NP -o Hi -t PGM -p SE -w 500
```

Paired-end Illumina 1.3 reads, annotated genome available/required for the reference genome Hi_86-028NP.fasta with a single strain (Hi_00345) for alignment and variant calling. No SNP/indel matrices required:

```
/home/user/bin/SPANDx/SPANDx.sh -r Hi_86-028NP -o Hi -a yes -v Haemophilus_influenzae_86_028NP_uid58093 -t Illumina_old -s Hi_00345
```

GWAS – generation of input data compatible with PLINK

Running PLINK for GWAS analysis across antibiotic-resistant strains vs. antibiotic-sensitive strains:

```
/home/user/bin/SPANDx/GeneratePlink.sh -r Hi_86-028NP -i inGroup_antibiotic_resistant.txt -o OutGroup_antibiotic_sensitive.txt
```

Running PLINK for GWAS analysis as above but changing locus presence-absence cut-offs for customised outputs (default is 0.9):

```
/home/user/bin/SPANDx/GeneratePlink.sh -r Hi_86-028NP -i inGroup_antibiotic_resistant.txt -o OutGroup_antibiotic_sensitive.txt -c 0.95
```

Running PLINK for GWAS analysis as above, but testing against annotated outputs (useful for instant identification of variants!):

```
/home/user/bin/SPANDx/GeneratePlink.sh -r Hi_86-028NP -i inGroup_antibiotic_resistant.txt -o OutGroup_antibiotic_sensitive.txt -c 0.95 -a yes -v Haemophilus_influenzae_86_028NP_uid58093
```

AUTHORS AND CITATION

SPANDx was developed by Dr. Derek Sarovich (@DerekSarovich) and Dr. Erin Price (@Dr_ErinPrice), Menzies School of Health Research, Darwin, NT 0810, Australia.

Any feedback you have regarding SPANDx is most welcome. If you find an error or bug, please contact Derek and Erin at mshr.bioinformatics@gmail.com. Please include a detailed description of the error you encountered, the operating system you used and what happened to cause the error. In addition, please send the appropriate log files with the description.

If you used SPANDx and found it useful, please cite it! ©

Sarovich DS and Price EP. 2014. SPANDx: a genomics pipeline for comparative analysis of large haploid whole genome re-sequencing datasets. *BMC Res. Notes* **7**:618.

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