Microbe-ID: An open source toolbox for microbial genotyping and species identification

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Microbe-ID: An open source toolbox for microbial genotyping and species identification

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Abstract

Development of tools to identify species, genotypes, or novel strains of invasive organisms is critical for monitoring emergence and implementing rapid response measures. Molecular markers, although critical to identifying species or genotypes, require bioinformatic tools for analysis. However, user-friendly analytical tools for fast identification are not readily available. To address this need, we created a web-based set of applications called Microbe-ID that allow for customizing a toolbox for rapid species identification and strain genotyping using any genetic markers of choice. Two components of Microbe-ID, named Sequence-ID and Genotype-ID, implement species and genotype identification, respectively. Sequence-ID allows identification of species by using BLAST to query sequences for any locus of interest against a custom reference sequence database. Genotype-ID allows placement of an unknown multilocus marker in either a minimum spanning network or dendrogram with bootstrap support from a user-created reference database. Microbe-ID can be used for identification of any organism based on nucleotide sequences or any molecular marker type and several examples are provided. We created a public website for demonstration purposes called Microbe-ID (www.microbe-id.org) and provided a working implementation for the genus Phytophthora (www.phytophthora-id.org). In Phytophthora-ID, the Sequence-ID application allows identification based on ITS or cox spacer sequences. Genotype-ID groups individuals into clonal lineages based on simple sequence repeat (SSR) markers for the two invasive plant pathogen species P. infestans and P. ramorum. All code is open source and available on github and CRAN. Instructions for installation and use are provided at https://github.com/grunwaldlab/Microbe-ID.

Background
Development of tools for identification of species, genotypes or strains is critical for monitoring emergence of invasive organisms such as *Phytophthora ramorum* causing sudden oak death (Grünwald et al. 2008), *Hymenoscyphus fraxineus* causing ash dieback (Gross et al. 2014), *Aphanomyces astaci* causing crayfish plague (Holdich et al. 2009), *Cryptococcus gattii* causing cryptococcosis and meningitis (Byrnes et al. 2010), or Methicillin-resistant *Staphylococcus aureus* causing invasive MRSA disease (Klevens et al. 2007). Molecular markers provide a rapid means for identification, but require various bioinformatics tools for identification of species and/or novel genotypes. In eukaryotes, sequences from the rRNA internal transcribed spacer (ITS) region and various mitochondrial DNA regions are used to delineate species (Coleman, 2003, 2007). ITS and mtDNA markers are now the most widely used markers in plants (Coleman, 2007, 2009), fungi (James et al., 2006), corals (Grajales et al., 2007), and oomycetes (Cooke et al., 2012; Robideau et al., 2011) and have been coined “DNA barcodes” because of their broad ability to distinguish species (Schoch et al., 2012). Classification of individuals using various molecular markers has recently increased. Multi-locus sequence types (MLST) are being widely used by researchers working with bacterial taxa to reveal the identity of samples by classification relative to known reference strains (Maiden et al., 2013). Other molecular markers or methods used to distinguish genotypes might include microsatellites (or simple sequence repeats) to identify strains and clonal lineages (Cooke et al., 2012; Ivors et al., 2006), DNA sequences for specific genic regions (Maiden et al., 2013), single nucleotide polymorphism (SNP) genotyping using reduced representation approaches (Grünwald et al. 2016) such as RAD-seq (Etter et al., 2011) or genotyping by sequencing (GBS, Elshire et al., 2011), or genome wide SNP genotyping (Huang et al., 2009).
In addition to the molecular methods developed, different types of online databases have been implemented to identify species using these molecular methods within groups of organisms. Examples of these databases are FungiDB for fungi and fungal-like organisms (Stajich et al., 2011), EuPathDB for eukaryotic organisms (Aurrecoechea et al., 2013), and the Phytophthora database which allows entries by experts in the Phytophthora community from different labs or countries for different species of the genus (Park et al., 2008). We previously reported on our development of a database for Phytophthora species and genotype identification using web tools to identify species using common barcodes, enabling the conjunction of modern laboratory techniques with highly curated databases for species identification (Grünwald et al. 2011).

Our objective here was to report the development of a toolbox for microbe identification (Microbe-ID) that can readily be customized for sequence based species identification (Sequence-ID) or molecular marker-based identification of genotypes (Genotype-ID) for any group of organisms. Our objectives were two-fold: 1) to implement Microbe-ID as a demonstration site that is customizable for any group of organisms and 2) to demonstrate a working implementation at Phytophthora-ID.org version 2.0 with significant updates from version 1.0 (Grünwald et al. 2011). Microbe-ID includes two modules, Sequence-ID and Microbe-ID. Sequence analysis is implemented based on use of a well characterized barcode region for the genus Phytophthora, but can be implemented to use any barcode sequence of interest. Genotype analysis can be implemented to use a variety of marker data types. To demonstrate the breadth of the developed tools and applicability to the diversity of microorganisms, the following three examples were included: codominant microsatellite data (SSR/Microsatellite) for the oomycete P. ramorum (Grünwald et al. 2009), concatenated Multi Locus Sequence Type (MLST) or individual locus sequences for the bacterium Clavibacter michiganensis subsp. michiganensis (Tancos et al. 2015, Supplementary Figure 1), and dominant Amplified Fragment Length Polymorphism (Binary...
(AFLP) data, Supplementary Figure 2) for the oomycete *Aphanomyces euteiches* (Grünwald and Hoheisel, 2006). Moreover, Genotype-ID can be expanded to include other marker systems including gene sequences for resistance to antibiotics or fungicides as well as presence/absence polymorphisms for effector genes or other adaptive loci. Two sequence databases were developed that help us demonstrate the utility of Microbe-ID. These databases are for the genus *Phytophthora* (*Phytophthora*-ID) with two sequence databases containing over 110 species that are mostly plant pathogens (Kroon et al., 2012), and two genotyping databases for populations of the potato late blight pathogen, *P. infestans*, and the sudden oak death pathogen, *P. ramorum* (Genotype-ID) (Grünwald et al. 2008; Kamoun et al. 2015). All of these tools are readily customizable and open source ([http://www.github.com/grunwaldlab/microbe-ID](http://www.github.com/grunwaldlab/microbe-ID)), provided as a demonstration site ([http://microbe-id.org/](http://microbe-id.org/)), and a working implementation of Sequence-ID and Genotype-ID that we use in our own work for the genus *Phytophthora* ([www.phytophthora-id.org](http://www.phytophthora-id.org)). Finally, a companion paper describes application of Microbe-ID for the implementation of a new website, Gall-ID, with novel tools for identification of gall-forming bacteria (Davis et al. 2016).

**Microbe-ID: a toolkit for web-based genotype and species identification**

**The Microbe-ID toolbox.** We developed a template website named Microbe-ID ([http://microbe-id.org/](http://microbe-id.org/)) with two separate modules, Sequence-ID and Genotype-ID, for sequence-based species identification and genotyping, respectively. The website is written using bootstrap ([http://getbootstrap.com/](http://getbootstrap.com/)), a HTML, CSS, and JS framework for developing responsive, mobile first projects on the web. Specific instructions, code, and resources necessary for implementation of Microbe-ID are provided on the github repository ([https://github.com/grunwaldlab/Microbe-](https://github.com/grunwaldlab/Microbe-)}
The server currently hosting Microbe-ID is running Centos Linux release 6.6, NCBI BLAST 2.2.28+ (Altschul et al., 1990), MAFFT version 7.221 (Katoh et al. 2002), and R version 3.1.2.

Below we describe specific tools required for customization of each module.

**Navigating and using Microbe-ID.** Navigating the demonstration site of Microbe-ID (http://Microbe-ID.org) the user will encounter a menu bar with links to Sequence-ID, Genotype-ID, and an about page. The home page has a general description of the functionality and components implemented in Microbe-ID, as well as links to the github site. Since input from the user is entered into forms as a query, each form implemented in Microbe-ID is encoded to check that the format of the data supplied by the user is supported. If the format is incorrect, the page will prompt the user with an error message, making the use of Microbe-ID more user-friendly.

**Sequence-ID.** We created a module called Sequence-ID that uses BLAST analysis of common sequence loci for species identification (Figure 1). Sequence-ID includes a PERL_CGI script that permits the communication between a user interface form (implemented in HTML5) and a BLAST database for the desired sequence data. The form recognizes the input in FASTA format and uses BLASTN to search for the most similar DNA sequences in the marker database. The PERL-CGI script receives the information of the HTML tabulated output from BLAST and displays a table of hits to the end user. Sequence-ID is customizable as the FASTA data file can readily be updated using the makeblastdb program of the BLAST suite. It can also similarly be implemented for BLASTP analysis of amino acid sequences.

In the Sequence-ID webpage, the user will find two main tabs: A “Blast” tab in which the web-app is contained, and a “Help” tab which contains a link to a “Site Help” prompt. This “Site Help” prompt shows an example of the FASTA sequence format recognized by Sequence-ID. The user can copy the FASTA sequence and paste it in the “Blast” tab text form to perform a BLAST search on the example query, or can download the database in FASTA format. After the
BLAST is performed, the web page will provide a table including the BLAST search results for the query of interest using the BLAST alignment format.

**Genotype-ID.** Genotype-ID is a web application designed to be user-friendly, and to facilitate the interaction with R using the set of tools listed in Table 1 (Figure 2). To develop the web application, we used the R package ‘shiny’ (Rstudio, 2013). Shiny facilitates the interactions between user, server, and R (Figure 2). The shiny web framework relies on reactive programming, which allows dynamic deployment of traditional R scripts in response to data input through a website console whereby results generated in R are subsequently pushed to the end user. Thus, ordinary R packages, which otherwise require familiarity with the language, can be deployed behind a user-friendly interface. Genotype-ID interacts principally with the R packages ‘poppr’ (Kamvar et al., 2014), `adegenet` (Jombart, 2008), `ape` (Paradis et al, 2004), and `pegas` (Paradis, 2010) (Table 1).

Genotype-ID has three different modules, which are shown as tabs on the website, each specific to different molecular markers: SSR/microsatellite data (SSR/Microsatellite data), multilocus genotype sequence types (MLST data), and AFLP, RFLP, SNP and other binary datasets (Binary (AFLP) data). The user supplies a query via the web framework that is read into R and queried against a curated dataset. Genetic distances (Table 2) are calculated between the query and genotypes in the reference database, with relationship of the query to the database presented either as a UPGMA or neighbor-joining dendrogram with bootstrap support or a minimum spanning network. These methods reconstruct relationships of the query relative to the genotypes found in the curated database. Each module has its own customization scheme to analyze each particular molecular marker type. The SSR data tool calculates Bruvo’s distance (Bruvo et al. 2004) to reconstruct the UPGMA or NJ dendrogram and minimum spanning networks. The MLST data tool permits the comparison of user-submitted gene sequences in
FASTA format. The MLST data tool uses MAFFT (Katoh et al. 2002) to align each query sequence to the corresponding curated gene database. It then concatenates all separate alignments, calculates the genetic distance for the alignment, and reconstructs distance dendrograms and minimum spanning networks. Lastly, the binary tool uses molecular markers such as AFLPs or RFLPs to reconstruct relationships. This tool uses binary data (coded as 1 and 0) and different genetic distances (Table 2) to reconstruct the UPGMA or NJ dendrogram and the minimum spanning network.

The Microbe-ID implementation. Microbe-ID contains implementations of each of the modules for different custom databases: the SSR implementation uses nine diploid SSR loci for the oomycete species P. ramorum (Figure 3); the MLST implementation uses eight multilocus sequence types for the bacterial species Clavibacter michiganensis subsp. michiganensis; and a binary implementation that uses 56 loci for AFLP data for the oomycete Aphanomyces euteiches. Each of the implementations of Genotype-ID contains collapsible instructions on how to format the query and a link to download a tabulated file with example queries formatted for use in Genotype-ID. The user can download the spreadsheet in order to edit, copy, and paste their custom queries into the “Data input” form. To make analyses that use a random seed repeatable, the user has the option to specify a seed number. In MLST-ID and Binary-ID, the user can select the genetic distance to be used in the analysis. After the user inputs the query and runs the web application, the web page will proceed to the Analysis section, where the user can choose between two different visualizations, a distance tree with bootstrap support values or a minimum spanning network. If the distance tree is selected, the user can change the tree algorithm (either neighbor joining or UPGMA) and number of bootstrap replicates. The user can also download results as a PDF or in NEWICK format. For the minimum spanning network, the user can adjust the grey
scale for edge distances and download results as a PDF file. Implementations for other genetic
data and different visualizations can be added.

**Phytophthora-ID implementation.** Tools provided in Microbe-ID were implemented in
http://Phytophthora-ID.org, a functional website for identifying samples from the genus
Phytophthora, a group of economically important plant pathogens in the stramenopile branch of
the tree of life (Kamoun et al. 2015). Phytophthora-ID version 1.0 (Grünewald et al. 2011) had a
BLAST script to identify samples using the ITS barcode. Phytophthora-ID version 2.0 was
substantially revised and upgraded from the first iteration and now includes a faster BLAST
search implemented in Sequence-ID and a new genotype identification system implemented in
Genotype-ID.

The current version of Phytophthora-ID contains a Sequence-ID module customized for
two molecular barcodes used for Phytophthora species identification (ITS and cox spacer). This
particular version of the sequence identification tool was created in PERL-CGI, and permits the
search of a FASTA sequence query against a curated database of Phytophthora species. The
PERL-CGI for Phytophthora-ID version 2.0 was also redesigned to run directly on the server to
make the web application more stable and faster. In contrast, in Phytophthora-ID 1.0, the web
application was designed as a communication wrapper to an external cluster, thus rendering
functionality dependent on the external server.

We implemented the Sequence-ID module to use sequences from two genetic regions: the
nuclear internal transcribed spacer (ITS) and mitochondrial cox spacer region spanning the cox1
and cox2 loci. Databases were created using sequences from published Phytophthora species
descriptions (Blair et al., 2008; Martin and Tooley, 2003) or those that belong to classical
Phytophthora species described at least 25 years ago and are still recognized as valid (Erwin and
Ribeiro, 1996).
For the ITS region, we gathered a total of 211 sequences representing 108 species. For the cox spacer region, we created a database of 150 sequences representing 106 species. Laboratory protocols for preparing samples for sequence analysis are available on the website and in a previous publication (Grünwald et al., 2011). Additionally, a file with the complete set of ITS sequences, cox spacer sequences, GenBank accession numbers, and *Phytophthora* spp. can be downloaded from the website. Documentation on updates to the databases is provided on the website.

We implemented two modules for genotype identification of two *Phytophthora* species: *P. infestans* that causes potato late blight, and *P. ramorum* that causes sudden oak death. *P. infestans* has more than 18 reported clonal lineages (Hu et al., 2012), while *P. ramorum* has 4 reported clonal lineages (Grünwald et al., 2009; Van Poucke et al., 2012). To establish a database with a wide representation of clonal lineages in each species, we obtained and prepared DNA samples from *P. infestans* and *P. ramorum* that were collected from various regions of the world. A total of 48 *P. ramorum* isolates representing the 4 reported clonal lineages (NA1, NA2, EU1, EU2) were genotyped at nine SSR loci (Cooke et al., 2012; Prospero et al., 2007; Vercauteren et al., 2010; Grünwald et al., 2009; Ivors et al., 2006). Similarly, 11 *P. infestans* clonal lineages that are dominant in the US (including US11, US12, US8, US20, US21, US23, US24, EU4, EU5, EU8, EU13) represented by a total population of 96 isolates were genotyped at 11 SSR loci, using protocols from Lees et al. (2006) and Li et al. (2013). We constructed SSR reference databases compatible with the poppr/adegenet R packages that provide dendrograms and minimum spanning networks for queries. Bootstrap support can be calculated for dendrograms. To test the datasets, we used two queries per dataset, one sample of NA1 and NA2 clonal lineages each for *P. ramorum* and one sample of clonal lineages US-23 and US-8 clonal lineages for *P. infestans*.
All samples grouped with the corresponding reference lineages with high support values (Figures 4 and 5), demonstrating functionality of the Microbe-ID tool to identify samples correctly.

Conclusions

We constructed a web framework that can use a wide array of molecular markers to rapidly determine identities of species and genotypes. This web framework is provided as open source code on github (https://github.com/grunwaldlab/Microbe-ID). We implemented a demonstration available at http://Microbe-ID.org. New implementations for any organisms require reference databases, scripts, and web pages including choices from the set of computational tools shown in Table 1. The Microbe-ID framework is currently implemented for two fully functional sites, Phytophthora-ID (http://Phytophthora-ID.org/) and Gall-ID (http://Gall-ID.cgrb.oregonstate.edu/; Davis et al. 2016).

Modularity of Microbe-ID permits implementation of a range of markers using curated databases of known species or genotypes to determine identity of specimens. Use of a custom BLAST database permits identification of any sample of a species of interest by using any sequence-based molecular marker. Note that the amount of time required for BLAST execution is dependent on size of the databases. For the Sequence-ID ITS region implementation for Phytophthora, each run was completed and loaded to the web-app in less than one second, with results shown as a BLAST output table.

Most innovative in this web framework is the Genotype-ID module, which permits the researcher to use, in a simple, web-based interface, any type of molecular marker with a corresponding and curated reference dataset. Use of a shiny server for strain identification is completely novel and modularity permits use of virtually any molecular marker suitable for
existing R packages. For genotyping, population genetic markers demonstrated include SSR, MLST, and AFLP/RFLP data, but can also be expanded to presence/absence of genes or alleles or any other genetic features of interest. Custom implementations must include a reference dataset and custom R scripts. Implementations can be customized for virtually any organism and any molecular marker that can be analyzed in R.

Development of tools for species identification has increased in recent years; but lack of modularity and complicated methodological procedures make the process of species identification tedious and challenging for researchers without sufficient skills in computational biology. We developed an easy-to-set-up web framework, which permits quick and flexible deployment of species identification and genotyping tools using sequence, microsatellite/SSR, AFLP/RFLP, or MLST data for any organism. Given use of the bootstrap html framework, webpages are light in terms of computer resources required and are thus usable on any device including smartphones and tablets, and function with any browser. These tools have been deployed and used successfully for the genus *Phytophthora* and gall-forming bacteria, providing examples of two curated websites (Grünwald et al. 2011; Davis et al. 2016).

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**Table 1.** Open source computational tools required to install and deploy Microbe-ID on a server.

- = reference not available; see website provided for information.

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Table 2. Genetic distances implemented in Microbe-ID.

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Figure Legends

**Figure 1.** Screen capture of the Sequence-ID user interface of Microbe-ID. The interface includes tabs for input of barcodes and a dropdown help link that contains laboratory protocols, examples data, and logs of each query dataset. The input frame has a control scheme that only permits data in FASTA format.

**Figure 2.** Diagram representing implementation of Genotype-ID, which is comprised of a user interface file (index.html) and a server file (server.R). Each file communicates with the R framework (via shiny) and user (via HTML5). On the user side (left side), user input is provided by copy/paste of a query and selects/specifies the desired application modifiers (seed number, genetic distance calculation). This information is subsequently received and processed by the server file, prompting the application to run in R. On the server side (right side) a database file (Marker DB), R packages, and functions are retrieved and executed. When the run is complete, the server file provides output to the user interface file and displayed on the app output.

**Figure 3.** Screen capture of the Genotype-ID user interface of Microbe-ID. Shown is the SSR-ID module for *P. ramorum*. Genotype-ID includes an Instructions link with an example Excel file that the user can modify, copy, and paste into the data input form. There is no limit to the number of queries submitted. In the Analysis section, tabs are provided to select either between a distance tree or a minimum spanning network rendering.

**Figure 4.** Results of SSR-ID for NA1 and NA2 queries of *P. ramorum* provided in the example data file. Each color represents a clonal lineage pre-assigned to each reference sample (NA1,
NA2, EU1, EU2) with queries colored in red. **A.** UPGMA tree with 1,000 bootstrap replicates and support values above branches. Queries are represented in red and all are correctly placed with reference samples of the presumptive clonal lineage while also representing the relationship between clonal lineages in the reference dataset. **B.** Minimum spanning network reconstruction. Edge shade and width are inversely proportional to Bruvo’s distance as shown in the horizontal scale bar. Queries are represented in red and placed in nodes with the most similar reference sample in the dataset, indicating the NA1 query is most similar to the PR-12-044 reference sample and the NA2 query is more closely related to the PR-05-156 and PR-12-103 samples, which also belong to the NA2 clonal lineage.

**Figure 5.** Results of SSR-ID queries for strains placed into the US8 and US23 clonal lineages of the potato late blight pathogen, *P. infestans*. Colors correspond to clonal lineages assigned to each reference sample (B, C, EU-13, EU-14, etc.) except for the queries which are colored in red. **A.** UPGMA tree with 1,000 bootstrap replicates with support values above branches. Queries are represented in red and all are correctly placed with samples of the presumptive clonal lineage while also representing relationships between clonal lineages in the reference dataset. **B.** Minimum spanning network reconstruction. Edge shade and width are proportional to Bruvo’s distance shown in the horizontal scale bar. Queries are represented in red nodes and appear in legend as ‘???’. Queries placed in nodes with the most similar reference sample, indicating that the US8 query is most similar to the PI-12-016 reference sample (US-8 clonal lineage) and the US23 query is most closely related to the PI-12-023 sample, part of the US-23 lineage.
Supplementary Figures

Supplementary Figure 1. UPGMA dendrogram of sub-module MLST-ID of Genotype-ID. The tree was constructed using 100 bootstrap replicates and MLST analysis of 8 genes (5 housekeeping genes and 3 virulence genes) of *Clavibacter michiganensis*. The query used is a concatenation of a WASH sample using all 8 genes. Queries are represented in blue. Note that all queries are correctly placed amongst samples of its presumptive clonal lineage while also representing relationships between lineages in the reference dataset.

Supplementary Figure 2. UPGMA dendrogram of sub-module Binary-ID of Genotype-ID using 100 bootstrap replicates for *Aphanomyces euteiches*. The queries used for this iteration of Binary-ID are two samples from a presumptive “Athena” origin. Queries are represented in red. Note that all queries were correctly placed amongst samples of its presumptive clonal lineage while also representing relationships between lineages in the reference dataset, indicating both queries are more closely related to the “Athena” population than the “Mt. Vernon” population.
Figure 1. Screen capture of the Sequence-ID user interface of Microbe-ID. The interface includes tabs for input of barcodes and a dropdown help link that contains laboratory protocols, examples data, and logs of each query dataset. The input frame has a control scheme that only permits data in FASTA format.
Figure 2. Diagram representing implementation of Genotype-ID, which is comprised of a user interface file (index.html) and a server file (server.R). Each file communicates with the R framework (via shiny) and user (via HTML5). On the user side (left side), user input is provided by copy/paste of a query and selects/specifies the desired application modifiers (seed number, genetic distance calculation). This information is subsequently received and processed by the server file, prompting the application to run in R. On the server side (right side) a database file (Marker DB), R packages, and functions are retrieved and executed. When the run is complete, the server file provides output to the user interface file and displayed on the app output.
Genotype-ID

Choose a tab to select the type of data and analysis.

SSR/Microsatellite data  MLST data  Binary (AFLP) data

SSR analysis for *P. ramorum*

Paste and submit microsatellite data (see reference files for format), and then choose a tab for a distance tree analysis or a minimum spanning network.

Click on the following links to extend the contents of the webpage:

*Instructions*  *Data Input*

Data Input

Random Seed (This will affect bootstrap values and the layout of the minimum spanning network.)

9449

Submit genotype

Analysis

Select between a distance tree with bootstrap support values or a minimum spanning network

Distance Tree with Bootstrap  Minimum Spanning Network

Figure 3. Screen capture of the Genotype-ID user interface of Microbe-ID. Shown is the SSR-ID module for *P. ramorum*. Genotype-ID includes an Instructions link with an example Excel file that the user can modify, copy, and paste into the data input form. There is no limit to the number of queries submitted. In the Analysis section, tabs are provided to select either between a distance tree or a minimum spanning network rendering.
Figure 4. Results of SSR-ID for NA1 and NA2 queries of P. ramorum provided in the example data file. Each color represents a clonal lineage pre-assigned to each reference sample (NA1, NA2, EU1, EU2) with queries colored in red. A. UPGMA tree with 1,000 bootstrap replicates and support values above branches. Queries are represented in red and all are correctly placed with reference samples of the presumptive clonal lineage while also representing the relationship between clonal lineages in the reference dataset. B. Minimum spanning network reconstruction. Edge shade and width are inversely proportional to Bruvo’s distance as shown in the horizontal scale bar. Queries are represented in red and placed in nodes with the most similar reference sample in the dataset, indicating the NA1 query is most similar to the PR-12-044 reference sample and the NA2 query is more closely related to the PR-05-156 and PR-12-103 samples, which also belong to the NA2 clonal lineage.
Figure 5. Results of SSR-ID queries for strains placed into the US8 and US23 clonal lineages of the potato late blight pathogen, *P. infestans*. Colors correspond to clonal lineages assigned to each reference sample (B, C, EU-13, EU-14, etc.) except for the queries which are colored in red. A. UPGMA tree with 1,000 bootstrap replicates with support values above branches. Queries are represented in red and all are correctly placed with samples of the presumptive clonal lineage while also representing relationships between clonal lineages in the reference dataset. B. Minimum spanning network reconstruction. Edge shade and width are proportional to Bruvo’s distance shown in the horizontal scale bar. Queries are represented in red nodes and appear in legend as ‘??’. Queries placed in nodes with the most similar reference sample, indicating that the US8 query is most similar to the PI-12-016 reference sample (US-8 clonal lineage) and the US23 query is most closely related to the PI-12-023 sample, part of the US-23 lineage.