Introduction

Francisella tularensis is a highly virulent, facultative intracellular pathogen, which causes tularemia in humans and animals [1]. Transmission occurs via arthropod bites (mosquitoes and ticks), inhalation of contaminated dust, or ingestion of contaminated food or water. Tularemia is endemic in Sweden and has been reported since 1931 causing between 100–700 infections annually [2]. The most common clinical form is ulceroglandular tularemia and all reported human cases of tularemia in Sweden have been caused by F. tularensis subspecies holarctica (type B). The disease occurs with a seasonal pattern–being especially prevalent in summer and autumn. It is most common in Northern and Middle Sweden but during the last decade the infection has spread further south (SmirseNet2, Swedish National Surveillance System).

Recently, several molecular typing methods have been utilized in studies of genetic population structure of F. tularensis including: PFGE, whole genome microarrays, multi-locus variable-number of tandem repeat (VNTR) analysis (MLVA), insertion-deletion markers (INDELs), and single nucleotide polymorphism (SNP) [3–12]. SNPs are point mutations that are evolutionarily stable in clonally reproducing organisms such as F. tularensis [12]. This stability makes them useful for classifying bacterial populations into specific genetic groups where phylogenetic relationships among groups can be accurately inferred [13]. Several studies have constructed a global phylogenetic tree using canSNPs, subdividing the subspecies holarctica samples into 30 subpopulations found worldwide [6,8,11,12,14]. MLVA has been very useful for several bacterial species, which, like F. tularensis, have little genomic variation [15,16]. MLVA analysis using 25 VNTR markers on 139 F. tularensis subspecies holarctica samples from North America, Europe and Asia, revealed five subpopulations, which with one exception were found in several well separated geographic areas [7].

The aim of present study was to evaluate canSNPs, MLVA, and PmeI-PFGE for practical uses in epidemiological investigation of
Results

Molecular typing

To evaluate the effectiveness of SNP, MLVA and PFGE typing as methods for tracing source and place of infection within an endemic country, we analysed 127 clinical specimens (Table S1). We discovered that all three typing methods identified two major genetic groups with near-perfect agreement. Six samples were removed from canSNP analysis due to DNA quality issues. Analysis with canSNPs revealed that all *F. tularensis* samples were assigned to two major groups: B.7/8 (n = 24) and B.12 (n = 97) (Figure 1A). The proportion of B.12 phylogroup to B.7/8 was in agreement with previous publication [8]. MLVA analysis resulted in two major clusters, Tul-I and Tul-II that corresponded with the B.12 and B.7/8 lineages, respectively (Table S1; Figure 1A). Two samples falling within the B.12 lineage formed a distinct MLVA cluster, Tul-III (Table S1). The two major MLVA clusters (Tul-I and Tul-II) were based on differences in the two less discriminating markers, Ft-M22 and Ft-M24, with only two allelic variants each. Tul-III MLVA cluster differed from all other MLVA types in the allele combination of VNTR markers Ft-M24 and Ft-M22 with 1 and 4 copies, respectively. Two other canSNP groups (B.4 and B.10) that were not found in this study have been previously reported in Sweden [8,11,12]. However, samples of these phylogroups are rather rare in Sweden so their lack of presence in this study may be due to our smaller sample size.

PFGE analysis of the 124 clinical samples revealed three different but rather similar PFGE profiles, types 1–3 (Figure 2). Comparison of PFGE data to MLVA and canSNP data (Table S1) revealed that PFGE types 1 and 3 corresponded to the two major canSNP groups B.12 (Tul-I) and B.7/8 (Tul-II), respectively (Table S1). MLVA cluster Tul-III corresponded with PFGE type 2. Taken together, canSNP, MLVA, and PFGE were in general agreement when classifying samples into major groups.

Higher genetic resolution was obtained with SNP and MLVA analyses compared to PFGE. Samples within the B.12 SNP lineage were further classified into one of nine minor phylogroups (B.12/13, n = 11; B.14, n = 5; B.20/21/33, n = 13; B.21/22, n = 13; B.22, n = 22; B.23/14/25, n = 24, B.33/34, n = 6; B.34/35, n = 1; and B.35/36, n = 2) (Figure 1A; Table S1) using SNP-signatures previously described [6,11,12]. Combining all five VNTR markers resulted in 33 unique MLVA subtypes wherein 17 unique subtypes were the result of variation at the Ft-M3 locus, which was the most rapidly mutating marker. Tul-I samples were assigned into one of five smaller subgroups (A–E), with each alpha code genotype further divided into Ft-M3 subtypes. Tul-II samples were less diverse than Tul-I, resulting in 3 unique MLVA-types referred to as #8, #10, and #33. The number of alleles of the five markers varied exhibiting moderate to high diversity (D) ranging from 0.15–0.90 as previous described [7] (Table 1).

At this higher resolution level, MLVA and canSNP assignment of samples were not in consistent agreement. Samples within identical MLVA genotype did not fall into a single canSNP group, but rather fell into multiple SNP phylogroups (Figure 1A; Table S1). Several canSNP groups contained multiple MLVA genotypes (Figure 1A; Table S1). This lack of grouping agreement is due to the type of genetic markers (SNPs or VNTRs) targeted by each typing system. SNP mutations are highly stable and therefore are informative of the genetic relationships among groups of isolates [13]. MLVA typing is based on VNTR loci that can be prone to mutational instability and, therefore, often result in homoplasy, making inferences about relationships among groups of isolates unreliable. When SNP analysis and MLVA are used in a hierarchical scheme, accurate genetic relatedness among SNP-defined groups of *F. tularensis* samples can be confidently known (Figure 1A) [12,13] and the MLVA data provided finer levels of discrimination among samples within the same phylogroup.

Geographic distribution of SNP groups

A highly complex geographic pattern among the phylogroups exists at a national and regional scale in Sweden (Figure 1). Our data present a pattern of wide distribution of multiple phylogroups that are found in multiple relatively distant counties in Sweden. Samples from both major groups (B.7/8 and B.12) are represented in middle and northern Sweden, with the highest density found in middle Sweden. Southern Sweden is sparse and is represented by samples classified in closely related terminal phylogroups within B.12 lineage (B.33/34 and B.34/35). Nearly all phylogroups identified in this study are present in middle Sweden, making this region the center of diversity, as previously described [6].

As a consequence of wide distribution of multiple phylogroups, extensive phylogenetic diversity was found within individual counties in middle Sweden. The richest diversity is found in Stockholm area (Figure 1B, county AB), representing seven diverse phylogroups, each with multiple MLVA subtypes. Örebro, a city in middle Sweden, was co-localized with 3 distinct phylogroups and MLVA subtypes (Table S1 and Figure 1, county T). Most samples appeared to cluster along major waterways, such as rivers and coastal areas of the Swedish east coast, the Baltic Sea. There was no obvious geographical correlation to age and gender of the patients (data not shown), but adults were significantly more commonly infected than children. Water bodies, like the Dalälven river, appears to be linked to multiple phylogroups that are diverse (Figure 1B, county W).

In contrast, Ljusnan river (county X in Figure 1B) is linked to seventeen case-patient samples that are highly genetically similar despite the 12 year span (1995–2006). All samples, except one, belonged to subgroup B.22 MLVA D4. The remaining sample fell in the B.21/22 subgroup, which is the closest relative to B.22 samples. Despite this pattern, 3 B.22 samples were found in 3 other nearby counties in middle Sweden, including Stockholm (Figure 1B), which may reflect recent dispersal from the founding source. Taken together, small regional sections or water bodies are co-localized with multiple phylogroups with great genetic diversity.

Discussion

The complex phylogeographic pattern of tularemia distribution in Sweden poses a great challenge to accurately identify a source and place of infection for any given *F. tularensis* specimen within Sweden (Figure 1). Our attempts to correlate geographic origin of the samples to specific genetic types did not generate a clear cut result (Figure 1B) despite employing a canSNP typing scheme that provided higher genetic resolution [6] than in the study performed by Karlsson et al 2013 and colleagues. This SNP typing scheme allowed us to place 121 clinical specimens, collected throughout Sweden over a span of 16 years, on the existing global phylogenetic tree for *F. tularensis* subsp. *holaretica* (Figure 1A) [6]. All phylogroups identified in Sweden have also been found in distant nations abroad, except B.22, which is restricted to Sweden. Given this broader dispersal range, it is no surprise that samples within these specific phylogroups are broadly dispersed in Sweden across multiple counties and not restricted to a small geographic region. The broad distribution of closely related phylogroups is central to the argument that *F. tularensis* is a rapidly dispersed
Figure 1. Detailed phylogeographic patterns of 103 human patient samples from Sweden. A) Existing global canSNP phylogeny of *F. tularensis* subsp. *holarctica* (Gyuranecz et al) wherein subgroups are indicated as circles and reference strains as stars. Gray coloration indicates subgroups not identified in this study. Noted for each mapped subgroup are the MLVA genotypes and n values (i.e. number of strains). The country code for each sequence genome is indicated: HUN, RUS, SWE, USA. B) Geographic distribution of groups on separate maps of Sweden based on membership to one of the two major subgroups (*B.7/8* and *B.12*). County code is indicated on the map (left). Total number of samples per county is represented by the circle size. Multiple subgroups within a single county are represented proportionally on a pie chart comprised of the colors corresponding to subgroups.

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organism. The basis for this rapid dispersal remains unclear, but recent reports suggest dispersal could be facilitated by the migration of birds [19,20].

Despite the phylogeographic complexity, patterns emerge that suggest the possibility of identifying SNP markers that could be meaningfully linked to geographic regions or at least narrow the geographic range of possible places. An example of this is the SNP for B.22 phylogroup, which is comprised of samples found only in Sweden [6,11,12]. It is interesting to note that unlike all other phylogroups in this study, B.22 SNP was identified from a Swedish genome (FSC200) linked to the Ljusnan river in county X (Figure 1). This B.22 SNP was highly specific to FSC200 genome. All other phylogroups are defined by SNPs discovered from genomes found in other nations (USA, Russia, and Hungary) [6,11,12]. Most case-patients (16/17) linked to the Ljusnan river, collected over a 12 year time frame (1995-2006) (Figure 1B), typed as B.22 MLVA D4 (Table S1). The striking genetic similarity suggests that they all recently emerged from a common ancestor. The temporal pattern rules out a single outbreak season and provides insight into the stability of this genotype. Taken together, these data suggest that the B.22 D4 genotype originated from a common ecological niche that spanned the approximately 70 kilometres distance of the river. Intense localized sampling of beaver populations may verify this hypothesis. Beavers, which are common on this river, could be a viable source given that these semi-aquatic rodent species have shown evidence of seroconversion to *F. tularensis* [21].

Three B.22 samples were found in other neighboring counties, one B.22 D4 type was found in Stockholm and two B.22 samples with D25 & D32 subtypes were found in counties W and S (Figure 1B). The B.22 samples located in multiple counties may reflect very recent dispersal events, which is supported by the MLVA subtype data, or errors in epidemiological records.

The B.22 SNP provides a striking line of evidence that supports the pattern that closely genetically related samples tend to have closer geographic proximity despite the complex phylogeographic landscape found in Sweden. Extrapolating from this model, SNPs that are highly specific to the sequenced genome may be found most frequently among samples recovered from the same location where the reference genome originated. Such SNPs would be both relationally and geographically informative and, therefore, useful in identifying a likely place of infection. That said, due to the rapid dispersal of *F. tularensis*, the geographical attribution of the B.22 SNP may be time limited.

The results of the present study indicate that SNP typing schemes, designed from geographically informative SNPs, combined with a MLVA typing scheme have the potential to be used as a standalone typing method in outbreak investigations. However, since our attempts to correlate geographic origin of the samples to specific genetic types did not generate a clear cut result despite employing a canSNP typing scheme that provided high genetic resolution we conclude that whole-genome sequencing (WGS) would be the most appropriate tool for effectively linking specific genetic type to geography. Comparing the WGS of an unknown sample to WGS databases of archived Swedish strains

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**Table 1. VNTR markers.**

| VNTR marker | Repeat size (nt) | No. of repeats* | No. of alleles* | Diversity** | Inside orf*** |
|-------------|-----------------|-----------------|----------------|-------------|---------------|
|             | Min | Max |                  |                |               |
| Ft-M3       | 9   | 8   | 25              | 17            | 0.90          | Yes           |
| Ft-M6       | 21  | 4   | 6               | 3             | 0.61          | Yes           |
| Ft-M20      | 12  | 2   | 4               | 3             | 0.15          | Yes           |
| Ft-M22      | 6   | 3   | 4               | 2             | 0.34          | Yes           |
| Ft-M24      | 21  | 1   | 2               | 2             | 0.36          | No            |

*Data obtained in this study.

1The individual marker diversity (D) was calculated as D=1-\[\sum\text{(allele frequency)}^2\].

2Location within an open reading frame.

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maximizes the likelihood of revealing those rare geographically informative SNPs among genetic near matches. This insight may prove useful for future epidemiological investigation practices.

Materials and Methods

Clinical samples

The Public Health Agency of Sweden receives continuously clinical specimens for primary diagnostics of *F. tularensis* from physicians. All samples were received during 1994 to 2010 from patients of both genders, varying ages (1–89 years), and from diverse regions in Sweden (Table S1). Since tularemia is a noticeable disease in Sweden, cases had been reported to the Swedish National Surveillance System by the treating physician. If no likely place of exposure had been included, the patients were contacted. We extracted *F. tularensis* subspecies holarctica DNA from patients suffering from ulceroglandular tularemia: 3 clinical samples taken directly from the wound site and 124 cultured samples. Genomic DNA was prepared using two commercially available DNA extraction kits, the QIAamp tissue protocol (Qiagen, Stockholm, Sweden) for samples from 1994 to 2005 and the NucliSens magnetic extraction protocol (Biomerieux, Gothenburg, Sweden) for *F. tularensis* samples and wound specimens from 2006–2010.

canSNP

CanSNP analysis using 26 previously published assays was performed on the 127 clinical samples as described (Figure 1A) [6,11,12].

MLVA

The markers Ft-M3, Ft-M6, Ft-M20, Ft-M22 and Ft-M24 were amplified as described [7]. The forward primers (Invitrogen Life Technologies, Paisley, United Kingdom), Ft-M3 and Ft-M20 were fluorescent labelled with NED, Ft-M6 and Ft-M24 with 6-FAM, and HEX for Ft-M22 respectively. PCR was performed in 12.5 μl reaction mixture containing 10 mM Tris-HCl, pH 8.3, 50 mM KCl, 2.5 mM MgCl2, 0.5 U AmpliTaq DNA polymerase (Applied Biosystems, Stockholm, Sweden), 0.1 mM deoxyribonucleoside triphosphates, 0.6 μM of each primer with the addition of 0.5–1.0 ng template. The reaction mixture was incubated at 94°C for 5 minutes and then cycled at 94°C for 30 s, 58°C for 30 s, and 72°C for 30 s, and finally at 72°C for 5 minutes. Amplicons were diluted 30 times and pooled into two combinations a) Ft-M3, Ft-M6, and Ft-M22 and b) Ft-M20, and Ft-M24, respectively. 1 μl of each pool was analysed with capillary electrophoresis (3130 Genetic Analyzer, POP7-polymer, and GeneScan TM-500 ROX TM size standards, Applied Biosystems, Stockholm, Sweden). The GeneMapper (Applied Biosystems) software was used to determine the size of the amplicons and to calculate the number of repeats at each VNTR marker. The diversity (D) for each VNTR marker was calculated as D = 1-∑(allele frequency)² [17].

PFGE

PFGE analysis was performed on *F. tularensis* samples [18]. Agarose plugs were slices and incubated in 10 U of restriction enzyme *PmeI* (Biolabs, New England) for 3 hours at 37°C. Electrophoresis was performed in 1% agarose with a switch time of 1.79 to 10.71 s at 6 V/cm for 24 hours at 14°C. *Salmonella enterica* serotype Braenderup strain H9812 restricted with *XbaI* was used for gel normalization. Gels were stained with gel red and gel images captured by using a Gel Doc 1000 imager (Bio-Rad).

PFGE images were analysed using BioNumerics v 6.01 (Applied Maths, Sint-Martens-Latem, Belgium). Unique PFGE patterns were analysed and compared manually for band polymorphism.

Geomapping

To obtain phylogeographic patterns we mapped the phylogenetic groups on a geographic map of Sweden at the county level resolution (Figure 1B). Of the 127 patients, 24 were excluded from this analysis due to unknown or uncertain location of exposure or lack of genotype information. The study protocol was approved by the Regional Ethical Review Board in Stockholm (2008/020-31/2).

Supporting Information

Table S1  *F. tularensis* clinical specimens from Sweden included in this study.

(XLSX)

Author Contributions

Conceived and designed the experiments: TW DB SL PK. Wrote the paper: TW DB PK SL. Contributed reagents/materials/analysis tools: DB TW SL PK. Performed the experiments: TW IH CM DW DB. Analyzed the data: TW DB MH SL.

References

1. Ellis J, Oyston PC, Green M, Tither RW (2002) Tularemia. Clin Microbiol Rev 15: 631–646.
2. Payne L, Arneborn M, Tegnell A, Giesecke J (2003) Endemic tularemia, Sweden. Emerg Infect Dis 11: 1440–1442.
3. Broekhuizen M, Larsson P, Johannson A, Byström M, Eriksson U, et al. (2003) Genome-wide DNA microarray analysis of Francisella tularensis strains demonstrates extensive genetic conservation within the species but identifies regions that are unique to the highly virulent *F. tularensis* subsp. tularensis. J Clin Microbiol 41: 2924–2931.
4. Farlow J, Smith KL, Wong J, Abrams M, Lytle M, et al. (2001) Francisella tularensis. Emerg Infect Dis 11: 15: 631–646.
5. García Del Blanco N, Dobson ME, Vela AI, Le De La Puente VA, Gutierrez CR, et al. (2002) Genotyping of Francisella tularensis strains by pulsed-field gel electrophoresis, amplified fragment length polymorphism fingerprinting, and 16S rRNA gene sequencing. J Clin Microbiol 40: 2964–2972.
6. Gyuranecz M, Birdsell DN, Splettstoesser W, Seibold E, Beckstro¨m-Sternberg SM, et al. (2012) Phylogeography of Francisella tularensis subs. holarctica, Europe. Emerg Infect Dis 18: 290–293.
7. Johannson A, Farlow J, Larsson P, Dukerich M, Chambers E, et al. (2004) Worldwide genetic relationships among Francisella tularensis isolates determined by multiple-locus variable-number tandem repeat analysis. J Bacteriol 186: 5088–5098.
8. Karlsson E, Svensson K, Lindgren P, Byström M, Sjödin A, et al. (2013) The phylogeographic pattern of Francisella tularensis in Sweden indicates a Scandinavian origin of Euro-Asian tularemia. Environ Microbiol 15: 654–665.
9. Larsson P, Svensson K, Karlsson L, Guala D, Granberg M, et al. (2007) Canonical insertion-deletion markers for rapid DNA typing of Francisella tularensis. Emerg Infect Dis 13: 1725–1732.
10. Staples JE, Kubota KA, Chalcraft LG, Mead PS, Petersen JM (2006) Epidemiologic and molecular analysis of human tularemia, United States, 1964–2004. Emerg Infect Dis 12: 1113–18.
11. Svensson K, Back E, Eliasson H, Berglund L, Granberg M, et al. (2009) Landscape epidemiology of tularemia outbreaks in Sweden. Emerg Infect Dis 15: 1937–1947.
12. Vogler AJ, Birdsell DN, Price LB, Zhgenti E, Babuadze G, et al. (2009) Phylogeography of Francisella tularensis: Global expansion of a highly fit clone. J Bacteriol 191: 2474–2484.
13. Keim P, Van Ert MN, Pearson T, Vogler AJ, Huyuh LY, et al. (2004) Anfrax molecular epidemiology and forensics: using the appropriate marker for different evolutionary scales. Infect Genet Evol 4: 205–211.
14. Chanturia G, Birdsell DN, Kekelidze M, Zhgenti E, Babuadze G, et al. (2011) Phylogeography of Francisella tularensis subspecies holarctica from the country of Georgia. BMC Microbiol 11: 159.
15. Keim P, Price LB, Klevytska AM, Smith KL, Schupp JM, et al. (2000) Multiple-locus variable-number tandem repeat analysis reveals genetic relationships within Bacillus anthracis. J Bacteriol 182: 2928–2936.
16. Klevytska AM, Price LB, Schupp JM, Worsham PL, Wong J, et al. (2001) Identification and characterization of variable-number tandem repeats in the Yersinia pestis genome. J Clin Microbiol 39: 3179–3185.
17. Weir BS (1990) Genetic data analysis: methods for discrete population genetic data analysis. Sinauer Associates, Inc., Sunderland, Mass.
18. Swaminathan B, Barrett TJ, Hunter SB, Tauxe RV, CDC PulseNet Task Force (2001) PulseNet: the molecular subtyping network for foodborne bacterial disease surveillance, United States. Emerg Infect Dis 7: 382–389.
19. Carvalho DD, Klein S, Akkus Z, ten Kate G, Schinkel A et al. (2012) Estimating 3D lumen centerlines of carotid arteries in free-hand acquisition ultrasound. Int J Comput Assist Radiol Surg 7: 207–215.
20. Padeski PI, Ivanov IN, Popov B, Kantardjiev TV (2010) The role of birds in dissemination of Francisella tularensis: first direct molecular evidence for bird-to-human transmission. Epidemiol Infect 138: 376–379.
21. Morner T, Sandstedt K (1983) A serological survey of antibodies against Francisella tularensis in some Swedish mammals. Nord Vet Med 35: 82–85.