Populations of Stored Product Mite *Tyrophagus putrescentiae* Differ in Their Bacterial Communities

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**Background:** *Tyrophagus putrescentiae* colonizes different human-related habitats and feeds on various post-harvest foods. The microbiota acquired by these mites can influence the nutritional plasticity in different populations. We compared the bacterial communities of five populations of *T. putrescentiae* and one mixed population of *T. putrescentiae* and *T. fanetzhangorum* collected from different habitats.

**Material:** The bacterial communities of the six mite populations from different habitats and diets were compared by Sanger sequencing of cloned 16S rRNA obtained from amplification with universal eubacterial primers and using bacterial taxon-specific primers on the samples of adults/juveniles or eggs. Microscopic techniques were used to localize bacteria in food boli and mite bodies. The morphological determination of the mite populations was confirmed by analyses of CO1 and ITS fragment genes.

**Results:** The following symbiotic bacteria were found in compared mite populations: *Wolbachia* (two populations), *Cardinium* (five populations), *Bartonella*-like (five populations), *Blattabacterium*-like symbiont (three populations), and *Solitalea*-like (six populations). From 35 identified OTUs\textsuperscript{97}, only *Solitalea* was identified in all populations. The next most frequent and abundant sequences were *Bacillus*, *Moraxella*, *Staphylococcus*, *Kocuria*, and *Microbacterium*. We suggest that some bacterial species may occasionally be ingested with food. The bacteriocytes were observed in some individuals in all mite populations. Bacteria were not visualized in food boli by staining, but bacteria were found by histological means in ovaria of *Wolbachia*-infested populations.

**Conclusion:** The presence of *Blattabacterium*-like, *Cardinium*, *Wolbachia*, and *Solitalea*-like in the eggs of *T. putrescentiae* indicates mother to offspring (vertical) transmission. Results of this study indicate that diet and habitats influence not only the ingested bacteria but also the symbiotic bacteria of *T. putrescentiae*.

**Keywords:** *Tyrophagus putrescentiae*, bacteria, symbiont, feeding, *Blattabacterium*, *Wolbachia*, 16S rRNA
INTRODUCTION

Mites, like insects, derive nutritive advantages from persistent associations with microorganisms (Van Asselt, 1999; Dillon and Douglas, 2004; Douglas, 2015). Microorganisms associated with insects can synthesize various nutrients, provide essential amino acids and contribute to digestive processes (Dillon and Dillon, 2004; Douglas, 2009). Domestic mites are inhabitants of human-related habitats such as homes, carpets, beds, and stored food (Spiekasma, 1997). Although these mites are commensals on the trophic level, due to allergen production and vectoring their microorganisms are of medical importance (Colloff, 2009). It is hypothesized that mites invaded human-related environments through two different routes: house dust mites (HDMs; e.g., *Dermatophagoides pteronyssinus* and *D. farinae*) likely shifted from a parasitic lifestyle back to a commensal life style (Klimov and OConnor, 2013), while ancestors of stored product mites (SPMs; e.g., *Acarus siro*, *Lepidoglyphus destructor*, *Tyrophagus putrescentiae*) were free-living and invaded human houses via the nests of birds and small mammals (OConnor, 1979, 1982).

*Tyrophagus putrescentiae* (Schrank, 1781) has been reported from agricultural soils (Smrz and Jungova, 1989), commercial bumblebee colonies (Rozej et al., 2012), and the nests of birds and small mammals (Solarz et al., 1999). This mite is very common in human-created habitats such as dust in urban environments, medical and laboratory facilities, farms, the food industry (Franz et al., 1997; Solarz et al., 2007), and in fungal and insect cultures in laboratories (Duck et al., 2001). The most typical food sources of *Tyrophagus putrescentiae* are protein and fat-rich substances such as grain germ, nuts, sunflower oil, rape seeds, cheese, ham, and dry dog food (Zakhvatkin, 1959; Robertson, 1961; Hughes, 1976; Garcia, 2004; Palvyos et al., 2008; Erban et al., 2015, 2016; Rybanska et al., 2015). Our broad hypothesis in work reported here is that successful colonization of such a wide range of habitats is facilitated by an inherently broad food plasticity in *Tyrophagus putrescentiae* that is likely due to symbiotic microbes.

In general, the ancestors of stored-product mites were fungivorous (OConnor, 1979, 1982). Fungivory is well-documented for *Tyrophagus putrescentiae* (Smrz and Catska, 1987; Hubert et al., 2004; Nesvorna et al., 2012). The mite can nutritionally benefit from interactions with bacteria or both fungi and bacteria. *Tyrophagus putrescentiae* produces bacteriolytic enzymes which hydrolyze the cell walls of gram positive bacteria (Erban and Hubert, 2008). *Tyrophagus putrescentiae* has been found to host bacterial communities in the gut, parenchymal tissues and reproductive tract with various interactions (Hubert et al., 2012a; Kopecky et al., 2014a,b). *Tyrophagus putrescentiae* is associated with bacterial parasites or symbionts (*Cardinium* and *Wolbachia*) inhabiting the reproductive tract and fat body (Kopecky et al., 2013; Brown and Lloyd, 2015). Another association was described for neutral and alkaline proteases, and exo-chitinase producing *Bacillus cereus* in *Tyrophagus putrescentiae*. *B. cereus* was presented in all samples and the exoenzymes can interact to utilization of the food sources for *Tyrophagus putrescentiae*; however, the addition of *B. cereus* to the diet led to a substantial suppression of mite population growth (Erban et al., 2016). In contrast, another bacterium, *Micrococcus lysodeikticus*, had no significant influence on the population growth of *T. putrescentiae* (Erban and Hubert, 2008). However, those associated bacteria affect the fitness of mites; therefore, their indirect effects on habitat colonization or diet utilization by mites are expected.

The above facts suggest that interactions of mites with microorganisms are important for adaptation to a nutritional food source in human-made habitats as well as in soil. A diet switches from plant-derived food to various fungal species caused changes in the bacterial community associated with the gut and parenchymal tissues of *Tyrophagus putrescentiae* (Smrz, 2003; Hubert et al., 2012b) and induced bacteriocytes (sometimes called as extraintestinal bacterial bodies or bacteriome; Smrz and Catska, 1989; Smrz, 2003; Smrz and Soukalova, 2008). It is likely that *Tyrophagus putrescentiae* acquires bacteria with chitinolytic activity (i.e., *B. cereus* and *Serratia marcescens*) to digest chitin from fungal cell walls or mite bodies (Smrz et al., 1991; Smrz and Catska, 2010; Erban et al., 2016). The bacteria producing exo-chitinases can contribute to degradation of mite exuviae, mite bodies or food boluses consisting of chitin, including the peritrophic membrane (Erban et al., 2016).

In laboratory experiments adding antibiotics to the diet did not eliminate the bacteria in *Tyrophagus putrescentiae*; *Kocuria* and *Bacillus* were still present in surface cleaned mite body homogenates (Kopecky et al., 2014b). Our recent study showed a population-specific density-dependent growth of *Tyrophagus putrescentiae* (Rybanska et al., 2015) and also indicated the possibility that *Tyrophagus putrescentiae* populations may differ in acquired bacteria (Smrz and Catska, 2010), suggesting a possible habitat influence on mite internal bacteria.

Here, we compare bacterial community of six distinct populations of *Tyrophagus putrescentiae*. The comparison is based on Sanger sequencing of bacterial 16S rRNA genes from adults/juveniles. Because we found diverse symbiotic bacterial community in these populations, we also focused on identification of these symbiotic bacteria in the eggs by bacterial tax specific primers. Finally, we compared sex ratios and guanine contents in *Tyrophagus putrescentiae* populations in relation to the presence of symbiotic bacteria. Limitations associated with the small amount of starting wild material do not allow reliable intra-population comparison of bacterial communities because prolonged mite culture maintenance under standardized laboratory conditions can change bacterial communities.

MATERIALS AND METHODS

Mites

The bacterial communities of three field and three laboratory populations of *Tyrophagus putrescentiae* (Schrank, 1781) were compared (Table 1). Morphological determination of populations was performed by under a compound microscope using characters described in Klimov and OConnor (2009,a,b). In addition, molecular markers were used for subsequent characterization of the populations. All these *Tyrophagus putrescentiae* populations were placed into IWAKI 25 cm² surface area
TABLE 1 | Sampled populations of *Tyrophagus putrescentiae* and their habitats at the time of collection.

| Abbreviations | Population | Habitat | Food          | Collected |
|---------------|------------|---------|---------------|-----------|
| Ham           | Ham        | Field   | Ham           | 2013      |
| Kop           | Koppert    | Laboratory | Grain-derived | 2012      |
| Lab           | Laboratory | Laboratory | Grain-derived | 1996      |
| Dog           | Dog        | Field   | Dry dog food  | 2007      |
| Phi           | Philips    | Laboratory | Dry dog food  | 2014      |
| Zvo           | Zvoleneves | Field   | Grain-derived | 2011      |

The year indicates since the colony was maintained in the Crop Research Institute, Prague, Czech Republic.

Sample Homogenization and DNA Extraction

Prior to DNA extraction, ethanol was replaced, in sequence, by bleach, and then washed three times with sterile phosphate-buffered saline (PBST: 3.2 mM Na$_2$HPO$_4$, 0.5 mM KH$_2$PO$_4$, 1.3 mM KCl, and 135 mM NaCl) with 0.05% w/w Tween® 20 detergent (Cat No. P9416, Sigma-Aldrich, St. Louis, MO, USA) to remove surface microflora. Samples with a total volume of 100 µL PBST were homogenized in a Radnoti tissue grinder (Cat. No. 440613; Monrovia, CA, USA). DNA from the homogenates was extracted using a Wizard® SV Genomic DNA Purification System (Cat No. A2361, Promega) according to the manufacturer's instructions. Extracted DNA was stored in a freezer at −20°C until the analyses were performed.

Molecular Markers for Identification of *T. putrescentiae* Populations

The primers (Table 2) spanning a portion of the 5.8S ribosomal DNA (rDNA), the full-length ITS2 region, and a part of the 28S rRNA and primers for the central part of the CO1 region of *T. putrescentiae* were used to confirm species determinations for mites used in this research (Yang et al., 2011). Amplifications were performed in a C1000 Thermal Cycler (Bio-Rad, Hercules, CA, USA). A total volume of 25 µl polymerase chain reaction (PCR) reaction mixture contained 200 µM dNTPs, 3 mM MgCl$_2$, forward and reverse primers (100 nM each), 0.5 unit Taq polymerase (all from Promega), and 5–30 ng of template DNA that included mite genomic DNA (for reaction conditions, see Yang et al., 2011). The amplicons were done for every sample. The resulting PCR products were visualized by agarose gel electrophoresis. Because we used samples at the population level, amplicons were purified with Wizard® SV Gel and the PCR product clean-up system Kit (Cat No. A9281, Promega) and cloned using pGEM®-T Easy Vector (Cat No. A1380 Promega). We usually selected four clones per sample, i.e., 12 clones per population. The obtained sequences were assembled with CodonCode Aligner, version 5.1.5 (CodonCode Corporation, Dedham, MA, USA).

Description of the Bacterial Community

Polymerase chain reaction amplification of 16S rRNA gene fragments was used to characterize the bacterial community...
using universal 27F/1492R primers (Barbieri et al., 2001). The PCR was done for every sample with the exception of laboratory population of *T. putrescentiae*. For the latter population, PCR conditions were same as was described above with the exception of annealing temperature (Table 2). The PCR products were cloned using the same protocol as above and 12 colonies were usually selected per sample (i.e., 30 per population) and sequenced in Macrogen. The same laboratory population of *T. putrescentiae* was analyzed by the same design on the same diet in our previous studies (Hubert et al., 2012a,b; Kopecky et al., 2014b) and we used the clones to characterize the bacterial community of *T. putrescentiae* laboratory population.

The presence of *Cardinium*, *Wolbachia*, Bartonella-like bacteria, Solitalea-like bacteria and Blattabacterium-like bacteria was determined by using taxon-specific primers (see Table 2). As a positive control, we used DNA samples of mites where bacteria had previously been identified using the same methodology (Kopecky et al., 2013, 2014a). The negative control was the double-distilled water used for the PCR master mix preparation. The reaction conditions were the same as described for the molecular markers used for identification of *T. putrescentiae* populations; the amplification conditions are specified in Table 2. All samples were tested in triplicate. PCR product of the expected size was considered as positive sample. One positive sample was considered as positive presence of bacteria in the population. A negative sample was a sample with no product detected by specific primers. The primer specificity was checked by sequencing of the randomly selected amplicons according to protocol described above.

### 16S rRNA Sequences

The 16S rRNA sequences were obtained by cloning of amplicons of universal eubacterial primers 27F/1492R (Table 2). The obtained sequences were assembled with CodonCode Aligner and the chimeras were removed using the Mallard and Pintail software (Ashelford et al., 2005, 2006). Altogether, 262 sequences obtained in this study and 176 sequences from previous studies (Hubert et al., 2012a,b; Kopecky et al., 2014b). The sequences were analyzed in MOTHUR v.1.36.0 software (Schloss et al., 2009). The sequences were aligned to Silva reference database (Quast et al., 2013), filtered and clustered and then analyzed and assigned to the operational taxonomic units (OTUs) defined at a distance level of 0.03. The sequences were identified using the Ribosomal Database Project's (RDP) naive Bayesian rRNA classifier as training set No. 14 (Wang et al., 2007). The representative sequences for individual OTUs27 were compared to the sequences in GenBank using nucleotide Blastn (Altschul et al., 1997).

### Phylogenetic Analysis

We conducted phylogenetic analyses on sequences of 16S rRNA, ITS-2 and CO1 of Flavobacteriales symbionts and *Wolbachia*. The references sequences originated from GenBank or RDP. Alignments of partial 16S rRNA gene sequences were performed using the SILVA Incremental Aligner v.1.2.11 (Pruesse et al., 2012). For the analysis of phylogenetic relationships, the best-fit model of nucleotide substitution was selected using jModelTest v.2.1.7 (Guindon and Gascuel, 2003; Darriba et al., 2012). Based on the selection, the general time reversible model with a proportion of invariable sites and gamma distribution (GTR+G+I), was employed to infer phylogenies in a Bayesian

### Table 2 | Primers used for characterization of *Tyrophagus putrescentiae* populations and detection of bacterial community.

| Specificity          | Target     | Name | Primer 5′-3′ | Tm (°C) | Length (bp) | Reference |
|----------------------|------------|------|--------------|---------|-------------|-----------|
| *T. putrescentiae*   | ITS2 region| 28S-F| CGACTTTTCAGAACGCAATTCG | 55      | 488         | Yang et al., 2011 |
|                      | 28S-R      | TACGGYTAACCTTGTTAGACTT | 50      | 1460       | Lane, 1991 |
| Bacteria             | CO1 region | 28S-R| AQAAGTTGTACCTGGCTCAG  | 50      | 377         | O’Neill et al., 1992 |
|                      | 28S-F      | R1492| TTAATGAACCATATATAGATGTC  | 50      | 1460       | Lane, 1991 |
| Wolbachia            | WpF        | TTTAGGCTGCTATGTTATAC  | 52      | 900        | Kopecky et al., 2014a |
| Bartonella-like      | Bart 1F    | 16S rRNA | TGCAGTGCAATCTACTTACAC  | 52      | 900        | Kopecky et al., 2014a |
|                      | Bart 2R    | 16S rRNA | TTGCTTGCGACGGCGAGCCCT  | 52      | 900        | Kopecky et al., 2014a |

Erban et al.  
Bacteria of *Tyrophagus putrescentiae*  

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framework in PhyloBayes-MPI, v.1.4e (Lartillot et al., 2009) and the maximum likelihood framework in PhyML v.3.0 (Guindon et al., 2010). Phylograms were visualized in MEGA 6 (Tamura et al., 2007).

Microanatomical Analyses

Sections
Mites were fixed in modified Bouin-Dubosque fluid (Smrz, 1989) for 3 days and then transferred to paraffin. The fixation fluid was replaced by 100% isopropyl alcohol for 12 h (two times), isopropyl alcohol/methyl benzoate (1/1 v/v) for 12 h (two times), methyl benzoate for 12 h (two times), benzene for 2 h, benzene/paraffin (1/1 v/v) for 12 h at 48°C, and paraffin for 12 h (two times) at 56°C (Hubert et al., 1999). Mites were transferred from containers to Peel-A-Way® embedding molds (Polysciences, Eppelheim, Germany) and embedded in Paraplast Plus® (Cat No. 39602004, Leica Biosystems, Nussloch, Germany) at 56°C. Paraffin blocks were sectioned to 4–6 µm sections on Microm HM 200 ErgoStar Microtome (Carl Zeiss, Jena, Germany).

Staining
The slides with sections were placed into different staining solutions equally. The sections were stained by Masson’s triple stain combined with PAS (periodic acid and Schiff agent) and Mann-Dominic and Ziehl–Neelsen staining for bacterial visualization.

Observations
At least 15 specimens per population were observed using an Axioskop compound microscope equipped with a digital camera and Axiovision software (Carl Zeiss).

Visualization in Microscopic Slides
The mites were mounted on permanent slides in Liquide de-Swan medium (distilled water, 20 mL; gum arabic, 15 g; chloral hydrate 50 g and glucose 3 g, glacial acetic acid 5 mL; Kramar, 1953). The sexing of adults and a quantitative estimate of guanine granules were performed using a compound microscope. Due to the transparency of the mite body, the granules were visible without staining (Figure 9). At least 100 specimens per one-sex determination and 30 per guanine quantification were observed. The semiquantitative categories were as follows: (0) no guanine granules, (1) low number, the granules filled less than 25% of the hysterosoma, (2) intermediate – granules filled between 25 and 50% of the hysterosoma, and (3) massive, granules filled more than 50% of the hysterosoma (Figure 9).

Statistical Analyses
The similarity of bacterial community in populations was based on analyses of 16S rRNA clones’ library. The shared file was constructed in MOTHUR. The diversity indexes, rarefaction and the comparison of population using Principal coordinate analyses with Euclidian and Jaccard distances were calculated in PAST 3 software (Hammer et al., 2001). The heatmap was constructed in XLSTAT (Addinsoft, New York, NY, USA) using filtering by standard deviation and reduction of low abundant

**FIGURE 1** | Phylogenetic analysis of the CO1 clones of *Tyrophagus putrescentiae* with the reference sequences from other related astigmatid mites (Webster et al., 2004; Dermauw et al., 2009; Klimov and O’Connor, 2009a,b; Bochkov et al., 2014; Ge et al., 2014; Sun et al., 2014). The phylogeny was inferred by Bayesian analysis of 46 partial CO1 gene sequences from observed populations, i.e., L, laboratory; K, Koppert; P , Phillips; D, Dog; H, Ham; and Z, field Zvoleneves. Branch lengths correspond to the mean posterior estimates of evolutionary distances (scale bar: 0.5). Branch labels indicate the Bayesian posterior probabilities and bootstrap support values from maximum likelihood analysis.
FIGURE 2 | Phylogenetic analysis of ITS clones of *T. putrescentiae* with the reference sequences from other related astigmatid mites (Noge et al., 2005; Liu et al., 2006; Klimov and O'Connor, 2008, 2013; Yang et al., 2011; Beroiz et al., 2014). The phylogeny was inferred by Bayesian analysis of 46 partial ITS gene sequences from the observed populations, i.e., L, laboratory; K, Koppert; P, Phillips; D, Dog; H, Ham; and Z, field Zvoleneves. Branch lengths correspond to the mean posterior estimates of evolutionary distances (scale bar: 0.1). Branch labels indicate Bayesian posterior probabilities and supporting bootstrap values from maximum likelihood analysis.
FIGURE 3 | Phylogenetic analysis of the Blattabacterium-like symbiont clones obtained from the Dog population of T. putrescentiae. The phylogeny was inferred by a Bayesian analysis of 24 partial 16S rRNA gene sequences with the reference sequences from RDP, i.e., 126 sequences of Blattabacterium, C. Brownia, and C. Sulcia, and the sequences of 398 type strains representing the order Flavobacteriales. Branch lengths correspond to the mean posterior estimates of evolutionary distances (scale bar: 0.5). Branch labels indicate Bayesian posterior probabilities and bootstrap support values from maximum-likelihood analysis. The phylogram was rooted using Bacteroides fragilis sequence NR074784 as an outgroup.

RESULTS

The Comparison of CO1 and ITS in T. putrescentiae Populations

The taxonomic comparison based on Bayesian analyses of CO1 showed that observed T. putrescentiae populations (i.e., Laboratory, Dog, Koppert, and Phillips) clustered together with the known CO1 sequences from GenBank. The exception occurred with the field populations from Zvoleneves, which formed two separate clusters: (i) four sequences (Z 1, 2, 4, 5) formed one cluster with A. siro and (ii) the next sequences clustered to Tyrophagus similis (Figure 1). Using a diagnostic CO1 dataset of 25 identified species of Tyrophagus (PBK, unpublished), we matched these four sequences with Tyrophagus fanetzhangorum. Seven other sequences clustered with T. putrescentiae, based on both GenBank data and our Tyrophagus dataset. The Bayesian analyses of ITS confirm the previous classification for the laboratory populations, Dog, Koppert, and Phillips with T. putrescentiae and similar to the CO1 data, the field Zvoleneves population (eight sequences) clustered with T. fanetzhangorum (Figure 2).

Flavobacteriales Symbiont (Blattabacterium-Like)

The Blattabacterium symbiont was amplified using specific primers designed for identification of Blattabacterium in cockroaches. The symbiont was present in mite populations from dog food only as indicated by 16S rRNA sequences from eubacterial primers, as well as the amplicons of 16S rRNA fragments obtained by specific Blattabacterium primers. A Bayesian analysis of the obtained sequences showed that the sequences clustered as a monophyletic lineage outside Blattabacterium (Clark and Kambhampati, 2003), Candidatus Brownia rhizoecola (Gruwell et al., 2010), C. Uzinora diaspidicola (Gruwell et al., 2007), and C. Sulcia muelleri (Moran et al., 2005; Figure 3).

Wolbachia

The sequences of Wolbachia were found in the clones of amplicons obtained by the clones form Wolbachia specific primers (WpF/WpR in Table 2) amplicons, i.e., 9 and 10 from Dog and Phillips populations, respectively. A Bayesian analysis of the obtained sequences and sequences in RDP showed that the T. putrescentiae sequences formed separate cluster (Figure 4). Our analyses showed more clusters of the sequences. The analyses differentiated the sequences from the insects and nematodes, with a few exceptions. The sequences were closer to clusters of nematodes and aphids. The sister group was formed by Wolbachia sequences from nematode Radopholus similis (Haegeman et al., 2009) and mite Torotrogla cardueli (Prostigmata: Syringophilidae; Glowska et al., 2015).

Comparison of Bacterial Community Based on 16S rRNA Gene

Obtained 16S rRNA sequences were deposited in GenBank (Accession Numbers: KX022128–KX022390) and combined to 176 sequences of the laboratory T. putrescentiae population which are available in GenBank (Accession Numbers: JN236405–JN236431; JX001234–JX001344, KJ635082–KJ635148; Hubert et al., 2012a,b; Kopecky et al., 2014b). All the sequences originated from amplification and cloning the amplicons from universal eubacterial primers. Altogether 42 OTUs were distinguished (Supplementary Table S1). The diversity

OTUs. The data describing sex ratio and guanine contents of mite bodies were tested by a chi-square test with Bonferroni correction in STATISTIX 9 software (Analytical Software, Tallahassee, FL, USA).
of bacterial community was similar for all populations of *T. putrescentiae* with exception of Zvoleneves population (Figures 5A,B). The bacterial community of mite population was formed from known/suspected symbionts or parasites: Bartonella-like bacteria (OTU97 1), Wolbachia (OTU97 4), Cardinium (OTU97 6), Blattabacterium-like symbiont (OTU97 7), Solitalea-like bacteria (OTU97 8; Figure 5C). The next most frequent and abundant OTUs97 were Bacillus (OTU97 2), Moraxella (OTU97 3), Staphylococcus (OTU97 5), Kocuria (OTU97 9), and Microbacterium (OTU97 10; Supplementary Table S1).

The populations differed in observed bacterial community as indicated by the 16S rRNA library. Principal coordinate analyses using Euclidian data matrix showed that bacterial communities of Phillips, and Dog mite populations were similar, while Zvoleneves, Laboratory and Ham populations were formed from different bacteria (Figure 6A). The first principal axis explained 60% and the second axis explained 23% of variation in the dataset. When Jaccard data matrix was calculated, the bacterial community was similar in Zvoleneves, Dog and Phillips populations, while Laboratory and Ham populations were different (Figure 6B). The first axis explained 59% and second axis explained 29% of variation in the data set. Composition between populations is reported in Figure 6C.

To confirm the distribution of bacterial taxa obtained from the analyses of cloned sequences from eubacterial (F24/R1492) amplicons, the samples were analyzed by taxa specific primers. The presence/absence of selected taxa of symbiotic/parasitic bacteria was confirmed by specific primers in the samples of adults and in the eggs (Table 3).
FIGURE 5 | The diversity and composition of bacterial communities in T. putrescentiae populations based on Sanger sequencing of the 16S rRNA gene clones, the amplicons originated from eubacterial primers (F24/R1492): (A) rarefaction analyses, (B) inverse Simpson diversity index, (C) heat map. Abbreviations for the T. putrescentiae populations are listed in Table 1.

The highest number of OTUs was found in the Zvoleneves population, while the lowest was in the Phillips population. Generally, no OTU97 (i.e., sequences with 3% of dissimilarity) was shared by all the populations. But when taxa specific primers were used the presence of Solitalea was confirmed in all populations including the eggs. The exception was Phillips T. putrescentiae population, when the Solitalea was not detected in the eggs (Table 3). Bartonella (OTU97 1) had
the highest relative numbers in the Koppert and laboratory strains, but \textit{Bartonella} was present in the Phillips, Dog and Ham populations as well. The amplicon from specific primers confirms the \textit{Bartonella} in all populations except field Zvoleneves population. \textit{Bartonella} was not found in the eggs except of the Phillips \textit{T. putrescentiae} population. \textit{Wolbachia} (OTU\textsubscript{97} 4) had high relative numbers in the Dog and Phillips populations. The specific primers confirm \textit{Wolbachia} in adults/juveniles of Dog and Phillips populations. However, in the eggs, \textit{Wolbachia} was found only in Dog \textit{T. putrescentiae}
TABLE 3 | The presence or absence of amplicons of bacterial 16S rRNA generated with universal and taxon-specific primers in different populations of Tyrophagus putrescentiae.

| Taxa                        | Population |
|-----------------------------|------------|
|                             | Ham        | Kop        | Lab        | Dog         | Phi         | Zvo         |
|                             | A/J        | E          | A/J        | E          | A/J        | E          | A/J        | E          |
| Bacteria                    | +          | +          | +          | +          | +          | +          | +          | +          |
| Bartonella-like             | +          | –          | +          | –          | +          | –          | –          | +          |
| Cardinium                   | +          | +          | +          | +          | –          | –          | +          | +          |
| Wolbachia                   | –          | –          | –          | –          | +          | +          | –          | –          |
| Blattabacterium-like        | –          | +          | –          | –          | –          | +          | –          | –          |
| Solitariae-like             | +          | +          | +          | +          | –          | –          | +          | +          |
| Spiroplasma                 | –          | –          | –          | –          | –          | –          | –          | –          |

A/J – adults/juveniles of mites, E – eggs; PCR product presence or absence is denoted by plus (+) or minus (−) sign.

population (Table 3). Bacillus (OTU97 2) formed bacterial communities in the Laboratory and Koppert populations. Moraxella (OTU97 3) formed bacterial communities in the Ham population only and the Blattabacterium-like symbiont (OTU97 7) in the Dog population only. However, the specific primers confirm Blattabacterium-like symbiont in both adults/juveniles of T. putrescentiae Dog population and in the eggs of laboratory T. putrescentiae population (Table 3). Cardinium (OTU97 6) was found in the clones of 16S rRNA of Ham, Koppert and Laboratory T. putrescentiae population. However, the presence was confirmed in all populations by specific primers with exception of Dog population and in the majority of populations was detected in the eggs.

**Microanatomical Description of Associated Bacteria**

Bacteria were present on the mite surfaces, in bacteriocytes, reproductive tracts and salivary glands (Figure 7D). The bacteria were not identified in food boli. The observed food boli (Figure 7A) contained concentrated mucoid substances or unidentified food fragments (Figures 7B,C). However, the mites ingested bacteria randomly, as indicated by the presence of bacteria in the foregut. In the Phillips and Dog populations we found bacteria inside ovaria (Figure 7E).

The bacteriocytes (Figure 8A) were found in some adults in all observed populations. The bacteria were localized in fat tissues and were of various sizes covering up to one third of the histological sections (Figure 8C). The bacteria in the bacteriocytes were formed by spherical (Figures 8B,D,E) particles with different staining compared to the rest of tissues; we identified these particles as bacteria. In addition, rod-shaped particles were found in the bacteriocytes in the laboratory population of T. putrescentiae (Figure 8F).

The guanin granulae filled hysterosoma of mites with different intensity (Figure 9). The populations significantly differed (Chi-square = 119.37, P < 0.001) in the guanine contents in their hysterosoma (Figure 10A). The differences were between Phillips population with prevailing low guanine contents and the rest of populations with high guanine contents (Figure 9).

**Estimation of Sex Ratio**

Males were present in all studied populations. As estimated under a compound microscope using slide-mounted mites, proportions male/(male+female) differed among the T. putrescentiae populations (Chi-square = 72.63, P < 0.001) formed following groups: (i) proportion about 0.5: Laboratory, Koppert, and Ham; (ii) female prevailing populations: Phillips, Dog, and Zvoleneves populations (Figure 10B).

**DISCUSSION**

We found significant differences in bacterial communities among various T. putrescentiae populations using Sanger sequencing of cloned bacterial 16S rRNA amplicons. Because of the complex taxonomy of T. putrescentiae and a possibility of the presence of cryptic species, we inferred single-gene phylogenetic trees aimed at species identification and analysis of sequences available in GenBank. We sequenced two genes most commonly used for Tyrophagus, COI and ITS-2. Both generated phylogenetic trees showed the presence of two distinct species, T. putrescentiae and T. fanetzhangorum. T. putrescentiae was present in five populations except. Zvoleneves population was a mixture of T. fanetzhangorum and T. putrescentiae. These analyses also allowed classification of previously unidentified GenBank sequences (Figures 1 and 2). We did not conduct a combined analysis of the two genes because many sequences from GenBank are available for one but not for both of these genes. We suggest that in the case of Zvoleneves population might be the results of bacterial community influenced by an additional factor, that is, the presence of T. fanetzhangorum having its own unique bacterial community.

The differences in bacterial communities confirm the previous results: there is clear variation in bacterial communities among different populations of mites, and this variation may be due to environmental factors such as differences in the diet, micro-habitat or simply the geographic sources, as shown previously for Rhizoglyphus robini (Zindel et al., 2013) and Carpoglyphus lactis (Hubert et al., 2015). The bacterial community of T. putrescentiae was formed mainly by symbiotic or parasitic bacteria, and
FIGURE 7 | The images show: (A) total view of the digestive tract of *T. putrescentiae*; (B) total view of the ventriculus and post-colon with ingested food; arrows point to food boli; (C) detail view of the post-colon of specimens with food boli formed from mucoid substances and fragments of diet (arrow); (D) detail view of the salivary glands with stained bacterial cells (arrows); (E) detail view of oocyte with stained bacterial cells (arrows). Staining: (A, B) Masson’s triple stain, (C) Ziehl–Neelsen, (D, E) Mann Dominici; Scales: (A, B) 100 µm, (D, E) 25 µm. c, colon; ca, caecum; cm, cheliceral muscles; fb, food bolus; oc, oocyte; oe, esophagus; pc, post-colon; sg, synganglion; v, ventriculus.

contained previously identified bacteria including *Bartonella*-like, *Cardinium* (Kopecky et al., 2013, 2014b), and *Wolbachia* (Brown and Lloyd, 2015). A *Blattabacterium*-like symbiont and *Solitalea*-like bacteria were newly identified for *T. putrescentiae*. The presence of *Cardinium*, *Wolbachia*, *Solitalea*-like bacteria and *Blattabacterium*-like bacteria in the eggs of some of the compared *T. putrescentiae* populations indicate symbiotic or parasitic association between these bacteria and *T. putrescentiae*. 
It also indicates mother to offspring (vertical) transmission via eggs of these bacteria. In one case (Phillips population), we found Bartonella in the eggs; however, in four T. putrescentiae populations, Bartonella was not observed in the eggs, suggesting that Bartonella is not vertically transmitted.

The next group of identified bacteria, i.e., Bacillus, Moraxella, Staphylococcus, Kocuria, and Microbacterium could have originated from ingested diet (Hubert et al., 2012a,b; Kopecky et al., 2014a,b). We suggest that ingested bacteria serve as source of nutrients for the mites whether
FIGURE 9 | Guanine deposits in *T. putrescentiae* (mite are mounted on permanent microscopic slides): (A) no guanine granules; (B) a low number, the granules fill less than 25% of the hysterosoma; (C) intermediate, the granules fill 25–50% of the hysterosoma; (D) massive, the granules fill more than 50% of the hysterosoma, (E) detail of the previous image. Scales: (A–D) 100 µm, (E) 25 µm.

by direct food source (Erban and Hubert, 2008) and/or that they predigest the food with exo-enzymes (Erban et al., 2016).

The interesting finding is the presence of *Solitalea*-like bacteria, which we identified previously in cloned 16S rRNA sequences from a laboratory strain of the grain mite *A. siro* (Hubert et al., 2012a,b). However, one sequence of *Pedobacter* (Solitalea-like) was found in a Chinese *T. putrescentiae* population from edible fungi by Qu et al. (2015). *Solitalea*-like bacteria were detected in this study in all the *T. putrescentiae* populations. Recently, we detected *Solitalea*-like bacteria in the reproductive tract and parenchymal tissues of *A. siro* (Hubert et al., under review). The bacteria were also present in the eggs (Hubert et al., under review). Given these findings, we suggest
that, similarly to Cardinium, this Solitalea-like bacteria might be either symbiotic or parasitic.

Wolbachia are well-known bacteria in insects and mites (Glowska et al., 2015); however, only a few such studies are available for Astigmata. A recent study demonstrated that T. putrescentiae miticides feeding on Drosophila corpses, including Wolbachia-infected corpses, are possible vectors of Wolbachia (Brown and Lloyd, 2015). In this study, we found that the Wolbachia sequences from T. putrescentiae are more similar to Wolbachia in quill mites Torotrogla cardueli (Acari: Syringophilidae; Glowska et al., 2015), aphids (Augustinos et al., 2011), and nematodes (Haegeman et al., 2009). We also found that the Wolbachia sequences from Dog and Phillips T. putrescentiae populations clustered altogether suggesting their high similarity. Wolbachia manipulates sexual reproductions in mites, causing cytoplasmic incompatibility and feminization of males (Breeuwer and Jacobs, 1996). The results for sex ratio observed in both populations of the T. putrescentiae where we identified Wolbachia indicated that males were present. At the population level, T. putrescentiae is infested by at least two bacteria with the known ability to manipulate sexual reproduction in their hosts, i.e., Cardinium and Wolbachia. However, our sampling method did not provide information about the distribution of bacteria among individual mites. We localized the bacteria in oocytes and the bacteria were not observed in all individuals. This means that the infection spreads in a mite population with some dynamics. The described horizontal transfer of Wolbachia from Drosophila to T. putrescentiae (Brown and Lloyd, 2015) might suggest a possibility for mite-to-mite transfer also in our system. However, infection in other species of astigmatid mites in laboratory populations was not observed (Kopecky et al., 2014a), and the results obtained here using Wolbachia-specific primers were similar (see Table 2). In addition, in the populations infected with Wolbachia, a lower proportion of mite males was observed.

Blattabacterium is a member of Flavobacteriales (Bacteroidetes), which are obligate mutualistic endosymbiotic bacteria living in the fat bodies of cockroaches (Sabree et al., 2009). The association between cockroaches and this bacterium allows cockroaches to successfully subsist on nitrogen-poor diets and exploit nitrogenous waste (Sabree et al., 2009). Other related taxa are symbionts of sap-feeding insects, i.e., Hemiptera: Diaspididae, Coccidae, Coccoidea (Moran et al., 2005; Grunwell et al., 2007; Rosenblueth et al., 2012). Here, we found a related bacterium in the population found on dog food. Dog food is designed for carnivorous mammals and contains high fat and nitrogen, which makes it very suitable for the rapid growth and development of T. putrescentiae (Rybanska et al., 2015). Acquiring such a bacterium for a T. putrescentiae population that infests dry dog food seems to be beneficial to T. putrescentiae, but the exact nature of mite and bacterium interactions is unknown.

The Blattabacterium and Bartonella-relative taxa are known to influence nitrogen metabolisms of their hosts (van Borm et al., 2002; Sabree et al., 2009). Because we found such bacteria in T. putrescentiae, we focused on the comparison of guanine contents in specimens in the analyzed populations. We hypothesized that Blattabacterium-like bacteria reduce nitrogen waste in the parenchymal tissue of T. putrescentiae in the same way as suggested for cockroaches (Sabree et al., 2009). Guanine is the waste product of nitrogenous metabolisms (Levinson et al., 1991a). It is present in feces and has a kairomone function as was observed in the model species A. siro (Levinson et al., 1991b). T. putrescentiae has the ability to accumulate guanine in the parenchymal tissues (Smrz and Catska, 1989). It was documented that T. putrescentiae on nitrogenous-rich diets, such as some fungi, accumulate guanine in the fat tissues (Smrz and Catska, 1989; Smrz, 2003). Massive feeding on a fungal-type diet leads to an irreversible accumulation of guanine called “white body syndrome,” in which guanine crystals form in large numbers in fat tissues and suppress the internal organ functions of mites, leading to a damage (Smrz and Catska, 1989). But the categorization of guanine granules in mite bodies did not confirm this suggestion, because
low guanine contents were observed in Phillips not in Dog populations. The high guanine content in our populations of Tyrophagus putrescentiae is a possible result of nitrogen un-balanced diets when the mites are not able to eliminate nitrogenous waste (Smrz, 2003).

Bacteriocytes were formed by bacteria in mite specimens that consume a fungal diet (Smrz and Catska, 1989). It was suggested that bacteria can enter from the gut to the bacteriocytes in the fat tissues and participate in chitin digestion (Smrz and Trelova, 1995; Smrz, 2003). In this study, we found bacteriocytes in specimens from all populations, but bacteriocytes were not present in all specimens. We did not identify bacteria forming bacteriocytes, although only Solitalea was identified in all populations. However, previously we observed Solitalea-like bacteria in A. siro, and bacteriocytes were not present (Hubert et al., 2012a).

We hypothesize that the habitat and food plasticity exhibited by Tyrophagus putrescentiae is mediated by different bacterial communities associated with mite populations. Of the 35 identified OTUs, only Solitalea was shared by all populations, indicating that different populations of Tyrophagus putrescentiae differ in their bacterial communities. Previously, large changes in bacterial communities were observed after a diet switch in Tyrophagus putrescentiae. However, those differences were due to bacterial taxa ingested with the diet, i.e., Bacillaceae (Bacillus, Lysinibacillus, Oceanobacillus, and Virgibacillus) and Micrococcales (Kocuria and Brevibacterium; Hubert et al., 2012b; Kopecky et al., 2014b). This was also supported by observations of a field strain of Tyrophagus putrescentiae in laboratory experiments on fungal diets, when Alcaligenes faecalis, Agrobacterium sp., Serratia marcescens, and Achromobacter sp. were identified by plating and cultivation of bacteria obtained from the homogenates of Tyrophagus putrescentiae specimens that had consumed the fungi (Smrz et al., 1991; Smrz and Soukalova, 2008; Smrz and Catska, 2010). Here, we observed changes in Tyrophagus putrescentiae populations in the presence and infestation rates of Bacillus, Staphylococcus, Kocuria, and Moraxella as indicated by sequencing of cloned 16S rRNA sequences. Among these, Moraxella was associated with a mite population feeding on ham. Bacteria forming food bolii were not observed in the studied populations of Tyrophagus putrescentiae. One explanation is that these bacteria are ingested in low numbers along with fragments of fungal mycelium or plant debris.

From a practical point of view, mites are important pests, and mite-caused damage increases as the population density increases. It is well-known that different food sources influence enzyme physiology of mites resulting in differences in population growths (Smrz and Catska, 1987; Erban and Hubert, 2008, 2010; Erban et al., 2009, 2016; Nesvorna et al., 2012; Rybanska et al., 2015). The results of this study indicate that diet and habitats influence not only the ingested spectrum of bacteria but also the symbiotic and parasitic taxa. These two components of the bacterial community can affect both mite fitness and population growth, which result in variability in the growth of various populations, causing higher interspecies variability than variability among species (Bowman, 1984). Therefore, the results of this study are also important for understanding nutritional biology of mites.

**AUTHOR CONTRIBUTIONS**

JH, TE, PK, JS, and TP: Scientific writing; JK: bioinformatics; MN: experiments, molecular biology, JH, JS and TE: experimental design, PK: taxonomy and interpretation, JH, and JS: microanatomy.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.01046

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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