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(54) Title: COMPOSITIONS AND METHODS FOR DETECTING NUCLEIC ACID FROM MOLLICUTES

(57) Abstract: Compositions, reaction mixtures, kits and methods used in amplifying and detecting nucleic acids from various species of the class Mollicutes. In certain aspects and embodiments, particular regions of the 23S rRNA or gene encoding said rRNA have been identified as preferred targets for nucleic acid amplification reactions of a sample suspected containing at least one species of Mollicutes. Some oligomers comprise tag regions. Some oligomers comprise target closing regions, promoter sequences, and/or binding moieties, such as homopolymeric and heteropolymeric nucleotide sequences. Samples can be from any source suspected of containing a species of the class Mollicutes. Preferred sample sources include bioreactors, cell lines, cell culture wares and pharmaceutical manufacturing wares, to name a few.

COMPOSITIONS AND METHODS FOR DETECTING NUCLEIC ACID FROM MOLLICUTES

CROSS REFERENCE TO RELATED APPLICATIONS

[01]. This application claims priority to U.S. Provisional Application Serial Number 61/219,674 filed on June 23, 2009, which is incorporated herein by reference in its entirety.

FIELD OF THE INVENTION

[02]. The field of the invention relates to amplification of nucleic acid from species of bacterium of the class *Mollicutes*.

BACKGROUND

- [03]. Bacteria in the class *Mollicutes* are parasitic organisms commonly contaminating eukaryotic cell culture systems. Members of the *Mollicutes* are among the smallest and simplest of the prokaryotes. Because of their small size and lack of a rigid cell wall, *Mollicutes* pass easily through filters intended to remove bacterial contaminants. *Mollicutes* contamination of a cell culture is problematic because it can negatively impact cell growth, alter metabolism and lead to unsafe final products. *Mollicutes* compete with cultured cells for essential nutrients and produce toxins that can cause cell death, all of which can impact the quality and productivity of cell cultures. These organisms are therefore of significant concern to the biopharmaceutical industry, which is dependent on continuous cell culture for the production of drugs, vaccines, and other biologics.
- [04]. Mollicutes include species from the genera Mycoplasma, Acholeplasma and Spiroplasma.

 Mollicutes can originate from mammalian, avian, insect, plant or fish cells. These organisms also have limited biosynthetic capabilities, making them dependent on external sources for essential nutrients and cofactors. Many species of Mollicutes have therefore evolved to become intracellular parasites. Thus, contaminating Mollicutes are present extracellularly and intracellulary in a host-cell environment.
- [05]. Current culture methods for detection of *Mollicutes* are limited by the time required for the growth of these fastidious organisms. As a result, detection of *Mollicutes* using these methods can take 28 days or longer, a timeframe that is not compatible with today's fast pace of pharmaceutical manufacturing and distribution. Access to rapid, simple and relatively inexpensive methods for the detection of *Mollicutes* would enable the routine testing and proactive quality control of cell cultures, raw materials, equipment, fixtures and the like.
- [06]. Compared to cell-culture methods, nucleic acid amplification tests are much more rapid detection methods. However, there is currently not a satisfactory nucleic acid amplification test available for detecting *Mollicutes* contamination of a cell-culture. Currently available nucleic acid amplification tests suffer from problems with identifying more than one or a few species of *Mollicutes* per assay. Further, for

the assays that are designed to identify more than one species of *Mollicutes*, there are commonly problems with cross reactivity to related bacteria or nucleic acids that may be present in the sample. Further still, the currently available tests fail to detect some of the *Mollicutes* species known to contaminate and negatively impact cell cultures. Thus, there is a need in the art for an improved nucleic acid amplification test for identifying *Mollicutes* contamination.

SUMMARY

- [07]. The present invention relates to compositions, reaction mixtures, kits and methods used in amplifying nucleic acids from various species of the class Mollicutes. Amplification reactions are uniplex reactions or, more preferably, are multiplex reactions. Amplified nucleic acids are then useful for a subsequent analysis. Subsequent analysis includes, amongst other types of analysis, nucleic acid detection, probe-based nucleic acid detection and real-time probe-based nucleic acid detection. In certain aspects and embodiments, particular regions of the 23S rRNA or gene encoding said rRNA have been identified as preferred targets for nucleic acid amplification reactions of a sample suspected containing at least one species of Mollicutes. Some oligomers comprise tag regions. Some oligomers comprise target closing regions, also referred to as tag-closing regions. Target closing regions have a nucleotide sequence that is substantially complementary to all or a portion of the target hybridizing region on the tagged amplification oligomer, thus allowing under certain conditions for the target closing region and the target hybridizing region to intramolecularly hybridize. The tagged oligomer is then considered to be in the closed/unavailable configuration. Some oligomers comprise promoter sequences. Some oligomers comprise binding moieties, such as homopolymeric and heteropolymeric nucleotide sequences. Compositions of the current invention comprise one or more separate oligomers useful in the amplification and detection of at least one species of Mollicutes. Samples can be from any source suspected of containing a species of the class Mollicutes. Preferred sample sources include bioreactors, cell lines, cell culture wares and pharmaceutical manufacturing wares, to name a few.
- [08]. In one embodiment of the invention, there is provided a composition for determining the presence or absence of one or more *Mycoplasma* species in a sample, said composition comprising a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1.
- [09]. In one aspect, said target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide

or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region is from 20 to 24 nucleobases in length. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:38. In another aspect said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:39.

- [10]. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof.
- [11]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:40. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:17. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:17.
- [12]. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer to allow for an intramolecular hybridization between the two regions under certain conditions. Preferably, the conditions are those that allow for the tagged oligomer to be in the closed position, while still allowing for steps of the reaction to be performed. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [13]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:5, SEQ ID NO:18 and SEQ ID NO:30. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:5. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:30.

In one embodiment, said composition further comprises an additional oligomer comprising a target [14]. hybridizing region containing a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5003 to 5045 of SEQ ID NO:1, wherein said additional oligomer is selected from the group consisting of a blocker oligomer, a primer oligomer and a promoter-based amplification oligomer. In one aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10. In one aspect, said additional oligomer is a promoter-based amplification oligomer. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification comprising a sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15.

- [15]. In one embodiment, said composition further comprises a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer or complement thereof. In one aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33.
- [16]. In one embodiment, said composition further comprises a first tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region

of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1, as is described above, and one or more additional tagged amplification oligomers. In one aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6, a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7 and a third additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:8.

[17]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer and the other of which is a blocker oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:5. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer

comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:5 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:5 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10.

- [18]. In one embodiment, said composition further comprises at least at least two additional oligomers, one of which is a tag-targeting oligomer, the other of which is a promoter based amplification oligomer. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13 and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15.
- In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1, as is described above; and the others of which can be selected from the group consisting of: an additional tagged amplification oligomer; a blocker oligomer; a tag-targeting oligomer; and a promoter based amplification oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In one aspect,

said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEO ID NO:5. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEO ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:5, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:5, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:7. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:8.

In one embodiment, there is a reaction mixture for generating from one or more Mycoplasma [20]. species in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1. In one aspect, said initial amplification product comprises a tag sequence that is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement

thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- [21]. In one embodiment, there is an amplification reaction mixture for generating from an initial nucleic acid amplification product containing a tag sequence or complement thereof, as described above, a further nucleic acid amplification product, wherein said amplification reaction mixture comprises a tagtargeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to specifically hybridize to said tag sequence, or complement thereof, that contained within the nucleotide sequence of said initial nucleic acid amplification product, and wherein said amplification reaction mixture comprises a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1.
- [22]. In one aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15. In another aspect, said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.
- [23]. In one aspect, there is provided an initial amplification product generated using a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1, wherein said initial amplification product contains a nucleotide sequence comprising a tag sequence region, and wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another

aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- [24]. In one aspect, there is provided an amplification product generated from said initial amplification product and containing a nucleotide sequence comprising a tag sequence region, wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said tag sequence region is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said tag sequence region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, a tag sequence region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- [25]. In one embodiment, there is provided a method for the *in vitro* amplification of a nucleic acid from one or more *Mycoplasma* species in a sample, comprising the steps of: (a.) contacting a sample with a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; and (b.) providing suitable conditions for performing an amplification reaction.
- [26]. In one aspect said nucleotide sequence contained within said tagged amplification oligomer target hybridizing region has at least one of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region of said tagged amplification oligomer is from 20 to 24 nucleobases in length. In another aspect, said target hybridizing region of said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:38. In another aspect, said target hybridizing region of said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:39.

[27]. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:17. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:17.

- [28]. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer to allow for an intramolecular hybridization between the two regions under certain conditions. Preferably, the conditions are those that allow for the tagged oligomer to be in the closed position, while still allowing for steps of the reaction to be performed. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [29]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:5, SEQ ID NO:18 and SEQ ID NO:30. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:5. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:30.
- [30]. In one aspect, said method further comprises the step of contacting said sample with an additional oligomer. In another aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10.

[31]. In one aspect, said method comprises contacting said sample with a tagged amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:5, a nucleotide sequence that is at least 95% identical to SEQ ID NO:17, a nucleotide sequence that is at least 95% identical to SEQ ID NO:18, a nucleotide sequence that is at least 95% identical to SEQ ID NO:30, a nucleotide sequence that is 100% identical to SEQ ID NO:17, a nucleotide sequence that is 100% identical to SEQ ID NO:18, a nucleotide sequence that is 100% identical to SEQ ID NO:30, a nucleotide sequence that is 100% identical to SEQ ID NO:40, or a nucleotide sequence that is 100% identical to SEQ ID NO:40, or a nucleotide sequence that is 100% identical to SEQ ID NO:41; and wherein said contacting step further comprises contacting said sample with a blocker oligomer. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said blocker comprises a sequence that is SEQ ID NO:10.

- In one aspect, said method at step (b.) results in an initial amplification product containing a tag [32]. sequence, or complement thereof. In another aspect, there is provided an initial amplification product with a nucleotide sequence comprising a tag sequence region, wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEO ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said initial amplification product has a 3' end defined by a blocker oligomer. In another aspect, said initial amplification product has a 3' end that is not defined by a blocker oligomer.
- [33]. In one embodiment of the current invention, there is provided a method for the *in vitro* amplification of an initial amplification product, comprising the steps of (c.) contacting the sample with at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect, the oligomers

of step (c.) and the oligomers of step (a.) are contacted to said sample at the same time point. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to the sample and a single amplification reaction is performed. In this aspect, steps (b.) and (d.) are combined to provide suitable conditions for performing an amplification reaction to generate an initial amplification product and for performing an amplification reaction generate a second amplification product therefrom. Though all of the amplification oligomers are present in this combined amplification reaction aspect, the tag-targeting amplification oligomers are configured to not participate in amplification until a tagged amplification oligomer is incorporated into an initial amplification product. In another aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at separate time points. In one aspect, the method comprises the step of (c.) providing to an amplification product generated in step (b.) a second reaction mixture comprising a tag-targeting amplification oligomer consisting essentially of a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer, and comprising a second amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction.

In one aspect for generating an amplification product, said sample is contacted with a first additional amplification oligomer that is a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said, or complement thereof in an initial amplification oligomer, and is contacted with a second additional amplification oligomer. In another aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:13. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33. In one aspect, said second amplification oligomer is a primer amplification oligomer, a promoter based amplification oligomer or a tagged amplification oligomer. In another aspect, said second amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to

SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:15.

- [35]. In one aspect for the *in vitro* amplification of a nucleic acid from one or more *Mycoplasma* species in a sample, there is provided at least one tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5; at least one blocker oligomer that is at least 95% identical to SEQ ID NO:10; at least one tag-targeting amplification oligomer that is at least 95% identical to SEQ ID NO:13 and at least one promoter-based amplification oligomer that is at least 95% identical to SEO ID NO:15. In another aspect, said at least one tagged amplification oligomer is 100% identical to SEQ ID NO:5. In another aspect, said at least one blocker oligomer is 100% identical to SEQ ID NO:10. In another aspect, said at least one tagtargeting amplification oligomer is 100% identical to SEQ ID NO:13. In another aspect, said at least one promoter-based amplification oligomer is 95% identical to SEQ ID NO:15. In another aspect there are provided one or more additional tagged amplification oligomers each or which could be selected from the group consisting of: a tagged amplification oligomer with at least 95% identity to SEO ID NO:6; a tagged amplification oligomer with at least 95% identity to SEQ ID NO:7, a tagged amplification oligomer with at least 95% identity to SEQ ID NO:8, a tagged amplification oligomer with 100% identity to SEO ID NO:6, a tagged amplification oligomer with 100% identity to SEQ ID NO:7 and a tagged amplification oligomer with 100% identity to SEO ID NO:8.
- [36]. In one embodiment there is further provided an analysis step, wherein an initial amplification product, an amplification product or both are analyzed. In one aspect, said analysis comprises a detection step to identify the presence or absence of an initial amplification product from an amplification reaction in step (b.), an amplification product generated in step (d.) or both initial amplification and amplification products generated in steps (b.) and (d.). In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is at least 95% identical to SEQ ID NO:14. In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is 100% identical to SEQ ID NO:14.
- [37]. In one embodiment of the invention, there is provided a composition for determining the presence or absence of one or more *Mycoplasma* species in a sample, said composition comprising a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2.

[38]. In one aspect, said target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:45. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:46. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:47. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:48. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4776 to 4798 of SEQ ID NO:2.

- [39]. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- [40]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:23. In another aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:49. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:23.
- [41]. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer to allow for an intramolecular hybridization between the two regions under certain conditions. Preferably, the conditions are those that allow for the tagged oligomer to be in the closed position, while still allowing for steps of the reaction to be performed. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [42]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a

sequence selected from the group consisting of: SEQ ID NO:6 and SEQ ID NO:24. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:6. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:24.

- [43]. In one embodiment, said composition further comprises an additional oligomer comprising a target hybridizing region containing a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4690 to 4732 of SEQ ID NO:2, wherein said additional oligomer is selected from the group consisting of a blocker oligomer, a primer oligomer and a promoter-based amplification oligomer. In one aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10. In one aspect, said additional oligomer is a promoter-based amplification oligomer. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4703 to 4732 of SEQ ID NO:2. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification comprising a sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15.
- [44]. In one embodiment, said composition further comprises a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer or complement thereof. In one aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33.

[45]. In one embodiment, said composition further comprises a first tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2, as is described above, and one or more additional tagged amplification oligomers. In one aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7 and a third additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:8.

[46]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer and the other of which is a blocker oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95%

identical to SEQ ID NO:10. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6.

- [47]. In one embodiment, said composition further comprises at least two additional oligomers, one of which is a tag-targeting oligomer, the other of which is a promoter based amplification oligomer. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15.
- [48]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2, as is described above; and the others of which can be selected from the group consisting of: an additional tagged amplification oligomer; a blocker oligomer; a tag-targeting oligomer; and a promoter based amplification oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional

NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEO ID NO:9, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:6, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:7. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:8.

[49]. In one embodiment, there is a reaction mixture for generating from one or more *Mycoplasma* species in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2, as is described above. In one aspect, said initial amplification product comprises a tag sequence that is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that

contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- [50]. In one embodiment, there is an amplification reaction mixture for generating from an initial nucleic acid amplification product containing a tag sequence or complement thereof, as described above, a further nucleic acid amplification product, wherein said amplification reaction mixture comprises a tagtargeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to specifically hybridize to said tag sequence, or complement thereof, that is contained within the nucleotide sequence of said initial nucleic acid amplification product, and wherein said amplification reaction mixture comprises a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4703 to 4732 of SEQ ID NO:2.
- [51]. In one aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15. In another aspect, said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.
- [52]. In one aspect, there is provided an initial amplification product that was generated using a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2, wherein said initial amplification product contains a nucleotide sequence comprising a tag sequence region, and wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases

in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- [53]. In one aspect, there is provided an amplification product generated from said initial amplification product and containing a nucleotide sequence comprising a tag sequence region, wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said tag sequence region is from 18 to 24 nucleobases in length. In another aspect, said tag sequence region is 19 nucleobases in length. In another aspect, said tag sequence region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- [54]. In one embodiment, there is provided a method for the *in vitro* amplification of a nucleic acid from one or more *Mycoplasma* species in a sample, comprising the steps of: (a.) contacting a sample with a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; and (b.) providing suitable conditions for performing an amplification reaction.
- [55]. In one aspect said nucleotide sequence contained within said tagged amplification oligomer target hybridizing region has at least one of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region of said tagged amplification oligomer is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region of said tagged amplification oligomer is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to

hybridize to a region of a target nucleic acid corresponding to residues 4776 to 4798 of SEQ ID NO:2. In another aspect, said target hybridizing region of said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:45, SEQ ID NO:46, SEQ ID NO:47 or SEQ ID NO:48.

- [56]. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; as sequence with 100% identity to SEQ ID NO:43 or complement thereof. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:23 or SEQ ID NO:49. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:23.
- [57]. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer to allow for an intramolecular hybridization between the two regions under certain conditions. Preferably, the conditions are those that allow for the tagged oligomer to be in the closed position, while still allowing for steps of the reaction to be performed. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [58]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:6 and SEQ ID NO:24. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:6. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:24.
- [59]. In one aspect, said method further comprises the step of contacting said sample with an additional oligomer. In another aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10.

[60]. In one aspect, said method comprises contacting said sample with a tagged amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:6, a nucleotide sequence that is at least 95% identical to SEQ ID NO:23, a nucleotide sequence that is at least 95% identical to SEQ ID NO:24, a nucleotide sequence that is 100% identical to SEQ ID NO:6, a nucleotide sequence that is 100% identical to SEQ ID NO:23, or a nucleotide sequence that is 100% identical to SEQ ID NO:24; and wherein said contacting step further comprises contacting said sample with a blocker oligomer. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said blocker comprises a sequence that is SEQ ID NO:9.

- [61]. In one aspect, said method at step (b.) results in an initial amplification product containing a tag sequence, or complement thereof. In another aspect, there is provided an initial amplification product with a nucleotide sequence comprising a tag sequence region, wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said initial amplification product has a 3' end defined by a blocker oligomer. In another aspect, said initial amplification product has a 3' end that is not defined by a blocker oligomer.
- [62]. In one embodiment of the current invention, there is provided a method for the *in vitro* amplification of an initial amplification product, comprising the steps of (c.) contacting the sample with at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at the same time point. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to the sample and a single amplification reaction is performed. In this aspect, steps (b.) and (d.) are combined to provide suitable

conditions for performing an amplification reaction to generate an initial amplification product and for performing an amplification reaction generate a second amplification product therefrom. Though all of the amplification oligomers are present in this combined amplification reaction aspect, the tag-targeting amplification oligomers are configured to not participate in amplification until a tagged amplification oligomer is incorporated into an initial amplification product. In another aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at separate time points. In one aspect, the method comprises the step of (c.) providing to an amplification product generated in step (b.) a second reaction mixture comprising a tag-targeting amplification oligomer consisting essentially of a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer, and comprising a second amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction.

[63]. In one aspect for generating an amplification product, said sample is contacted with a first additional amplification oligomer that is a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said, or complement thereof in an initial amplification oligomer, and is contacted with a second additional amplification oligomer. In another aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:13. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33. In one aspect, said second amplification oligomer is a primer amplification oligomer, a promoter based amplification oligomer or a tagged amplification oligomer. In another aspect, said second amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4703 to 4732 of SEQ ID NO:2. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that comprises a sequence that is at least 95% identical to SEO ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:15.

[64]. In one aspect for the *in vitro* amplification of a nucleic acid from one or more *Mycoplasma* species in a sample, there is provided at least one tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6; at least one blocker oligomer that is at least 95% identical to SEQ ID NO:9; at least one tag-targeting amplification oligomer that is at least 95% identical to SEQ ID NO:13 and at least one promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In another aspect, said at least one tagged amplification oligomer is 100% identical to SEQ ID NO:6. In another aspect, said at least one blocker oligomer is 100% identical to SEO ID NO:9. In another aspect, said at least one tagtargeting amplification oligomer is 100% identical to SEQ ID NO:13. In another aspect, said at least one promoter-based amplification oligomer is 95% identical to SEQ ID NO:15. In another aspect there are provided one or more additional tagged amplification oligomers each or which could be selected from the group consisting of: a tagged amplification oligomer with at least 95% identity to SEQ ID NO:5; a tagged amplification oligomer with at least 95% identity to SEQ ID NO:7, a tagged amplification oligomer with at least 95% identity to SEQ ID NO:8, a tagged amplification oligomer with 100% identity to SEQ ID NO:5, a tagged amplification oligomer with 100% identity to SEQ ID NO:7 and a tagged amplification oligomer with 100% identity to SEQ ID NO:8.

- [65]. In one embodiment there is further provided an analysis step, wherein an initial amplification product, an amplification product or both are analyzed. In one aspect, said analysis comprises a detection step to identify the presence or absence of an initial amplification product from an amplification reaction in step (b.), an amplification product generated in step (d.) or both initial amplification and amplification products generated in steps (b.) and (d.). In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is at least 95% identical to SEQ ID NO:14. In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is 100% identical to SEQ ID NO:14.
- [66]. In one embodiment of the invention, there is provided a composition for determining the presence or absence of one or more *Acholeplasma* species in a sample, said composition comprising a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 SEQ ID NO:3.
- [67]. In one aspect, said target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect,

said target hybridizing region is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:50. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:51. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:52. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:53. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:54. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:55. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1973 to 2006 of SEQ ID NO:3.

- [68]. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tagclosing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [69]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:7, SEQ ID NO:31, SEQ ID NO:56 or SEQ ID NO:57. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:7. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:31.
- [70]. In one embodiment, said composition further comprises an additional oligomer comprising a target hybridizing region containing a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1842 to 1884 of SEQ ID NO:3, wherein said additional oligomer is selected from the group consisting of a blocker oligomer, a primer oligomer and a promoter-based amplification oligomer. In one aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer that is at least 95% identical to SEQ

ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10. In one aspect, said additional oligomer is a promoter-based amplification oligomer. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1855 to 1884 SEQ ID NO:3. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification comprising a sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional oligomer is a promoter-based amplification oligomer is a promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15.

- [71]. In one embodiment, said composition further comprises a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer or complement thereof. In one aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33.
- [72]. In one embodiment, said composition further comprises a first tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3, as is described above, and one or more additional tagged amplification oligomers. In one aspect, said one or more additional tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said one or more additional

tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a third additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:8.

[73]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer and the other of which is a blocker oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:7. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:7 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:7 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:7 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10.

[74]. In one embodiment, said composition further comprises at least two additional oligomers, one of which is a tag-targeting oligomer, the other of which is a promoter based amplification oligomer. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15.

[75]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEO ID NO:3, as is described above; and the others of which can be selected from the group consisting of: an additional tagged amplification oligomer; a blocker oligomer; a tag-targeting oligomer; and a promoter based amplification oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEO ID NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:8. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEO ID NO:7. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEO ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical

to SEQ ID NO:7, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:7, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:8.

- [76]. In one embodiment, there is a reaction mixture for generating from one or more Acholeplasma species in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3, as is described above. In one aspect, said initial amplification product comprises a tag sequence that is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEO ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- [77]. In one embodiment, there is an amplification reaction mixture for generating from an initial nucleic acid amplification product containing a tag sequence or complement thereof, as described above, a further nucleic acid amplification product, wherein said amplification reaction mixture comprises a tag-

targeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to specifically hybridize to said tag sequence, or complement thereof, that contained within the nucleotide sequence of said initial nucleic acid amplification product, and wherein said amplification reaction mixture comprises a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1855 to 1884 of SEQ ID NO:3. In one aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15. In another aspect, said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.

[78]. In one aspect, there is provided an initial amplification product that was generated using a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEO ID NO:3, wherein said initial amplification product contains a nucleotide sequence comprising a tag sequence region, and wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEO ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEO ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, there is provided an amplification product generated from said initial amplification product and containing a nucleotide sequence comprising

a tag sequence region, wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said tag sequence region is from 18 to 24 nucleobases in length. In another aspect, said tag sequence region is 19 nucleobases in length. In another aspect, said tag sequence region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

[79]. In one embodiment, there is provided a method for the *in vitro* amplification of a nucleic acid from one or more *Acholeplasma* species in a sample, comprising the steps of: (a.) contacting a sample with a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and (b.) providing suitable conditions for performing an amplification reaction.

F801. In one aspect said nucleotide sequence contained within said tagged amplification oligomer target hybridizing region has at least one of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region of said tagged amplification oligomer is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region of said tagged amplification oligomer is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1973 to 2006 of SEQ ID NO:3. In another aspect, said target hybridizing region of said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:50, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:54 or SEQ ID NO:55. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEO ID NO:13. SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a

sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:7, SEQ ID NO:31, SEQ ID NO:56 or SEQ ID NO:57. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:7. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:31.

- [81]. In one aspect, said method further comprises the step of contacting said sample with an additional oligomer. In another aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:10. In one aspect, said method comprises contacting said sample with a tagged amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:7, a nucleotide sequence that is at least 95% identical to SEQ ID NO:31, a nucleotide sequence that is 100% identical to SEQ ID NO:31; and wherein said contacting step further comprises contacting said sample with a blocker oligomer. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said blocker comprises a sequence that is SEQ ID NO:10.
- [82]. In one aspect, said method at step (b.) results in an initial amplification product containing a tag sequence, or complement thereof. In another aspect, there is provided an initial amplification product with a nucleotide sequence comprising a tag sequence region, wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity

to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said initial amplification product has a 3' end defined by a blocker oligomer. In another aspect, said initial amplification product has a 3' end that is not defined by a blocker oligomer.

- [83]. In one embodiment of the current invention, there is provided a method for the in vitro amplification of an initial amplification product, comprising the steps of (c.) contacting the sample with at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at the same time point. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to the sample and a single amplification reaction is performed. In this aspect, steps (b.) and (d.) are combined to provide suitable conditions for performing an amplification reaction to generate an initial amplification product and for performing an amplification reaction generate a second amplification product therefrom. Though all of the amplification oligomers are present in this combined amplification reaction aspect, the tag-targeting amplification oligomers are configured to not participate in amplification until a tagged amplification oligomer is incorporated into an initial amplification product. In another aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at separate time points. In one aspect, the method comprises the step of (c.) providing to an amplification product generated in step (b.) a second reaction mixture comprising a tag-targeting amplification oligomer consisting essentially of a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer, and comprising a second amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction.
- [84]. In one aspect for generating an amplification product, said sample is contacted with a first additional amplification oligomer that is a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said, or complement thereof in an initial amplification oligomer, and is contacted with a second additional amplification oligomer. In another aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:13. In

another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33. In one aspect, said second amplification oligomer is a primer amplification oligomer, a promoter based amplification oligomer or a tagged amplification oligomer. In another aspect, said second amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1855 to 1884 of SEQ ID NO:3. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:15.

[85]. In one aspect for the *in vitro* amplification of a nucleic acid from one or more *Acholeplasma* species in a sample, there is provided at least one tagged amplification oligomer that is at least 95% identical to SEQ ID NO:10; at least one blocker oligomer that is at least 95% identical to SEQ ID NO:13 and at least one tag-targeting amplification oligomer that is at least 95% identical to SEQ ID NO:13 and at least one promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In another aspect, said at least one tagged amplification oligomer is 100% identical to SEQ ID NO:10. In another aspect, said at least one blocker oligomer is 100% identical to SEQ ID NO:13. In another aspect, said at least one tag-targeting amplification oligomer is 100% identical to SEQ ID NO:15. In another aspect there are provided one or more additional tagged amplification oligomers each or which could be selected from the group consisting of: a tagged amplification oligomer with at least 95% identity to SEQ ID NO:5; a tagged amplification oligomer with at least 95% identity to SEQ ID NO:6, a tagged amplification oligomer with 100% identity to SEQ ID NO:5, a tagged amplification oligomer with 100% identity to SEQ ID NO:6 and a tagged amplification oligomer with 100% identity to SEQ ID NO:6 and a tagged amplification oligomer with 100% identity to SEQ ID NO:6 and a tagged amplification oligomer with 100% identity to SEQ ID NO:6.

[86]. In one embodiment there is further provided an analysis step, wherein an initial amplification product, an amplification product or both are analyzed. In one aspect, said analysis comprises a detection step to identify the presence or absence of an initial amplification product from an amplification reaction in step (b.), an amplification product generated in step (d.) or both initial amplification and amplification products generated in steps (b.) and (d.). In another aspect, said detection step is a probe-based detection

step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is at least 95% identical to SEQ ID NO:14. In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is 100% identical to SEQ ID NO:14.

- [87]. In one embodiment of the invention, there is provided a composition for determining the presence or absence of one or more *Spiroplasma* species in a sample, said composition comprising a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.
- In one aspect, said target-hybridizing region of said tagged amplification oligomer may contain [88]. one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 or SEQ ID NO:62. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2017 of SEQ ID NO:4. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.
- [89]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:8, or SEQ ID NO:63. In another aspect, said tagged amplification oligomer comprises 100%

sequence identity to SEQ ID NO:8.

In one embodiment, said composition further comprises an additional oligomer comprising a target [90]. hybridizing region containing a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1903 to 1945 of SEQ ID NO:4, wherein said additional oligomer is selected from the group consisting of a blocker oligomer, a primer oligomer and a promoter-based amplification oligomer. In one aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:9. In one aspect, said additional oligomer is a promoter-based amplification oligomer. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1916 to 1945 SEQ ID NO:4. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a target hybridizing region comprising a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said additional oligomer is a promoter-based amplification comprising a sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional oligomer is a promoter-based amplification oligomer comprising a sequence that is 100% identical to SEO ID NO:15.

- [91]. In one embodiment, said composition further comprises a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer or complement thereof. In one aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33.
- [92]. In one embodiment, said composition further comprises a first tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15

to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, as is described above, and one or more additional tagged amplification oligomers. In one aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a tagged amplification oligomer comprising a nucleic acid sequence that is 100% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:6. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7. In another aspect, said one or more additional tagged amplification oligomers includes a first additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a second additional tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6 and a third additional tagged amplification oligomer that is at least 95% identical to SEO ID NO:7.

[93]. In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer and the other of which is a blocker oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:8. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8. In another aspect, said blocker oligomer comprises a

nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8 and said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10.

- [94]. In one embodiment, said composition further comprises at least two additional oligomers, one of which is a tag-targeting oligomer, the other of which is a promoter based amplification oligomer. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15.
- In one embodiment, said composition comprises at least two oligomers, one of which is a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, as is described above; and the others of which can be selected from the group consisting of: an additional tagged amplification oligomer; a blocker oligomer; a tag-targeting oligomer; and a promoter based amplification oligomer. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:8. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is at least 95% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification

oligomer comprises a nucleic acid sequence that is at least 95% identical to SEQ ID NO:7. In one aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10. In one aspect, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13. In another aspect, said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:9, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said tagged amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:8, said blocker oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:10, said tag-targeting amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:13, and said promoter-based amplification oligomer comprises a nucleotide sequence that is 100% identical to SEQ ID NO:15. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:5. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:6. In another aspect, said additional tagged amplification oligomer comprises a nucleic acid sequence that is 100% identical to SEQ ID NO:7.

[96]. In one embodiment, there is a reaction mixture for generating from one or more *Spiroplasma* species in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, as is described above. In one aspect, said initial amplification product comprises a tag sequence that is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a

nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- In one embodiment, there is an amplification reaction mixture for generating from an initial [97]. nucleic acid amplification product containing a tag sequence or complement thereof, as described above, a further nucleic acid amplification product, wherein said amplification reaction mixture comprises a tagtargeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to specifically hybridize to said tag sequence, or complement thereof, that contained within the nucleotide sequence of said initial nucleic acid amplification product, and wherein said amplification reaction mixture comprises a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1916 to 1945 of SEQ ID NO:4. In one aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15. In another aspect, said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.
- [98]. In one aspect, there is provided an initial amplification product that was generated using a tagged amplification oligomer comprising a target hybridizing region and a tag region, wherein said target hybridizing region is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, wherein said initial amplification product contains a nucleotide sequence comprising a tag sequence region, and wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and

SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, there is provided an amplification product generated from said initial amplification product and containing a nucleotide sequence comprising a tag sequence region, wherein the sequence of said tag region is from 10 to 25 nucleobases in length. In another aspect said tag sequence region is from 18 to 24 nucleobases in length. In another aspect, said tag sequence region is 19 nucleobases in length. In another aspect, said tag sequence region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag sequence region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

[99]. In one embodiment, there is provided a method for the *in vitro* amplification of a nucleic acid from one or more *Spiroplasma* species in a sample, comprising the steps of: (a.) contacting a sample with a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4; and (b.) providing suitable conditions for performing an amplification reaction.

[100]. In one aspect said nucleotide sequence contained within said tagged amplification oligomer target hybridizing region has at least one of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region of said tagged amplification oligomer is from 18 to 28 nucleobases in length. In another aspect, said target hybridizing region of said tagged amplification oligomer is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2017 of SEQ ID NO:4. In another aspect, said target hybridizing region of said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 or SEQ ID NO:62.

In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; as sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:8, or SEQ ID NO:63. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:8.

[101]. In one aspect, said method further comprises the step of contacting said sample with an additional oligomer. In another aspect, said additional oligomer is a blocker oligomer. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said additional oligomer is a blocker oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said additional oligomer is a blocker comprising a sequence that is SEQ ID NO:9. In one aspect, said method comprises contacting said sample with a tagged amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:8, a nucleotide sequence that is at least 95% identical to SEQ ID NO:63, or a nucleotide sequence that is 100% identical to SEQ ID NO:8; and wherein said contacting step further comprises contacting said sample with a blocker oligomer. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:10. In another aspect, said blocker comprises a sequence that is SEQ ID NO:9.

[102]. In one aspect, said method at step (b.) results in an initial amplification product containing a tag sequence, or complement thereof. In another aspect, there is provided an initial amplification product with a nucleotide sequence comprising a tag sequence region, wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag

sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said initial amplification product has a 3' end defined by a blocker oligomer. In another aspect, said initial amplification product has a 3' end that is not defined by a blocker oligomer.

[103]. In one embodiment of the current invention, there is provided a method for the in vitro amplification of an initial amplification product, comprising the steps of (c.) contacting the sample with at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at the same time point. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to the sample and a single amplification reaction is performed. In this aspect, steps (b.) and (d.) are combined to provide suitable conditions for performing an amplification reaction to generate an initial amplification product and for performing an amplification reaction generate a second amplification product therefrom. Though all of the amplification oligomers are present in this combined amplification reaction aspect, the tag-targeting amplification oligomers are configured to not participate in amplification until a tagged amplification oligomer is incorporated into an initial amplification product. In another aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at separate time points. In one aspect, the method comprises the step of (c.) providing to an amplification product generated in step (b.) a second reaction mixture comprising a tag-targeting amplification oligomer consisting essentially of a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer, and comprising a second amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect for generating an amplification product, said sample is contacted with a first additional amplification oligomer that is a tagtargeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said, or complement thereof in an initial amplification oligomer, and is contacted with a second additional amplification oligomer. In another

aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:13. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33. In one aspect, said second amplification oligomer is a primer amplification oligomer, a promoter based amplification oligomer or a tagged amplification oligomer. In another aspect, said second amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1916 to 1945 of SEQ ID NO:4. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said second additional amplification oligomer is a promoter-based amplification oligomer that comprises a sequence that is 100% identical to SEQ ID NO:15.

[104]. In one aspect for the *in vitro* amplification of a nucleic acid from one or more *Spiroplasma* species in a sample, there is provided at least one tagged amplification oligomer that is at least 95% identical to SEQ ID NO:9; at least one tag-targeting amplification oligomer that is at least 95% identical to SEQ ID NO:13 and at least one promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In another aspect, said at least one tagged amplification oligomer is 100% identical to SEQ ID NO:8. In another aspect, said at least one blocker oligomer is 100% identical to SEQ ID NO:9. In another aspect, said at least one taggeting amplification oligomer is 100% identical to SEQ ID NO:13. In another aspect, said at least one promoter-based amplification oligomer is 95% identical to SEQ ID NO:15. In another aspect there are provided one or more additional tagged amplification oligomers each or which could be selected from the group consisting of: a tagged amplification oligomer with at least 95% identity to SEQ ID NO:5; a tagged amplification oligomer with at least 95% identity to SEQ ID NO:6, a tagged amplification oligomer with at least 95% identity to SEQ ID NO:5, a tagged amplification oligomer with 100% identity to SEQ ID NO:5, a tagged amplification oligomer with 100% identity to SEQ ID NO:7.

[105]. In one embodiment there is further provided an analysis step, wherein an initial amplification product, an amplification product or both are analyzed. In one aspect, said analysis comprises a detection step to identify the presence or absence of an initial amplification product from an amplification reaction in step (b.), an amplification product generated in step (d.) or both initial amplification and amplification products generated in steps (b.) and (d.). In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is at least 95% identical to SEQ ID NO:14. In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is 100% identical to SEQ ID NO:14.

[106]. In one embodiment of the invention, there is provided a composition for determining the presence or absence of one or more species from the Mollicutes class in a sample, said composition comprising at least two tagged amplification oligomers, each of which individually comprises a target hybridizing region and a tag region, wherein individually, each of the target hybridizing regions of the at least two tagged amplification oligomers are selected from the group consisting of: a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.

[107]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 24 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1. In another aspect, said target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:38 or SEQ ID NO:39. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25

nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with 100% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[108]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:5, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:30, SEQ ID NO:40 and SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:5. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:17. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged

[109]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 18 to 28 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:45, SEQ ID NO:46, SEQ ID NO:47 or SEQ ID NO:48. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4776 to 4798 of SEQ ID NO:2. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a

sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[110]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:6, SEQ ID NO:23, SEQ ID NO:24 and SEQ ID NO:49. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:6. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:23. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:24.

[111]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 19 to 24 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:50, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:54 or SEQ ID NO:55. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1973 to 2006 of SEQ ID NO:3. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ

ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[112]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:7, SEQ ID NO:31, SEQ ID NO:56 or SEQ ID NO:57. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:7. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:31.

[113]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 25 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 or SEQ ID NO:62. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2017 of SEQ ID NO:4. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:8, or SEQ ID NO:63. In another aspect, said tagged amplification

oligomer comprises 100% sequence identity to SEQ ID NO:8.

[114]. In one aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:5 and a second tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:6 and a second tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:7 and a second tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:8 and a second tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6 or SEQ ID NO:7. In another aspect, said at least two tagged amplification oligomers comprises three tagged amplification oligomers each of the three individually being a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises four tagged amplification oligomers each of the four individually being a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8.

[115]. In one embodiment, said composition further comprises at least one additional oligomer, wherein said at least one additional oligomer is selected from the group consisting of: a blocker oligomer, a primer oligomer and a promoter-based amplification oligomer. In one aspect, one of said at least one additional oligomer is a blocker oligomer. In another aspect, said blocker oligomer is at least 95% identical to SEQ ID NO:9. In another aspect, said blocker oligomer is at least 95% identical to SEQ ID NO:10. In another aspect, said blocker oligomer is SEQ ID NO:9. In another aspect, said blocker oligomer is SEQ ID NO:9. In another aspect, said blocker oligomers, one of which is at least 95% identical to SEQ ID NO:9; or one of which is at least 95% identical to SEQ ID NO:10; or one of which is at least 95% identical to SEQ ID NO:9 and another of which is 100% identical to SEQ ID NO:10; or one of which is at least 95% identical to SEQ ID NO:10; or one of which is at least 95% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is 100% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is at least 95% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is at least 95% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is

100% identical to SEQ ID NO:10. In one aspect, one of said at least one additional oligomer is a promoter-based amplification oligomer. In another aspect, said promoter-based amplification oligomer comprises a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5003 to 5045 SEQ ID NO:1. In another aspect, said promoter-based amplification oligomer comprises a target hybridizing region that is at least 95% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprises a target hybridizing region that is 100% identical to SEQ ID NO:42. In another aspect, said a promoter-based amplification is at least 95% identical to SEQ ID NO:15. In another aspect, said promoter-based amplification oligomer is 100% identical to SEQ ID NO:15.

[116]. In one embodiment, said composition further comprises a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer or complement thereof. In one aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16.

[117]. In one embodiment, of the invention there is provided at least two tagged amplification oligomers selected from the group consisting of a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, as are described above; and there is at least one additional oligomer selected from the group consisting of: a blocker

oligomer; a tag-targeting oligomer; and a promoter based amplification oligomer. In one aspect, said blocker oligomer is at least 95% identical to SEQ ID NO:9 or SEQ ID NO:10. In another aspect, said blocker oligomer is 100% identical to SEQ ID NO:9 or SEQ ID NO:10. In one aspect, said tag-targeting amplification oligomer is at least 95% identical to SEQ ID NO:13. In another aspect, said tag-targeting amplification oligomer is 100% identical to SEQ ID NO:13. In one aspect, said promoter-based amplification oligomer is at least 95% identical to SEQ ID NO:15. In another aspect, said promoter-based amplification oligomer is 100% identical to SEQ ID NO:15. In one aspect, said composition comprises a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7, a blocker oligomer that is at least 95% identical to SEQ ID NO:9 and a promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In one aspect, said composition comprises a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, a blocker oligomer that is at least 95% identical to SEQ ID NO:10 and a promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In another aspect, said composition comprises a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, a blocker oligomer that is at least 95% identical to SEQ ID NO:9, a blocker oligomer that is at least 95% identical to SEQ ID NO:10 and a promoter-based amplification oligomer that is at least 95% identical to SEQ ID NO:15. In another aspect, said composition comprises SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10 and SEQ ID NO:15.

[118]. In one embodiment, there is a reaction mixture for generating from one or more species of the class *Mollicutes* in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises at least two tagged amplification oligomers, each of which individually comprises a target hybridizing region and a tag region, and wherein individually, each of the target hybridizing regions of the at least two tagged amplification oligomers are selected from the group consisting of: a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; a tagged amplification oligomer comprising a

of SEQ ID NO:3; and a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4, as are described above. In one aspect, said initial amplification product comprises a tag sequence that is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

[119]. In one embodiment, there is an amplification reaction mixture for generating from an initial nucleic acid amplification product containing a tag sequence or complement thereof, as described above, a further nucleic acid amplification product, wherein said amplification reaction mixture comprises a tagtargeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to specifically hybridize to said tag sequence, or complement thereof, that contained within the nucleotide sequence of said initial nucleic acid amplification product, and wherein said amplification reaction mixture comprises a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1. In one aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42. In another aspect, said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:42. In another aspect, said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15. In another aspect, said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.

[120]. In one embodiment, there is provided a method for the in vitro amplification of a nucleic acid in a sample, said nucleic acid being from one or more species in the class Mollicutes, comprising the steps of: (a.) contacting a sample with at least two tagged amplification oligomers, each of which individually comprises a target hybridizing region and a tag region, wherein said at least two tagged amplification oligomers are selected from the group consisting of: a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4; and (b.) providing suitable conditions for performing an amplification reaction.

[121]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 24 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1. In another aspect, said target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:38 or SEQ ID NO:39. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEO ID NO:13. SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is

substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[122]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a sequence selected from the group consisting of SEQ ID NO:5, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:30, SEQ ID NO:40 and SEQ ID NO:41. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:5. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:17. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:18. In another aspect, said tagged

[123]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 18 to 28 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEO ID NO:2. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEO ID NO:45, SEO ID NO:46. SEQ ID NO:47 or SEQ ID NO:48. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 4776 to 4798 of SEQ ID NO:2. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[124]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to a

sequence selected from the group consisting of SEQ ID NO:6, SEQ ID NO:23, SEQ ID NO:24 and SEQ ID NO:49. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:6. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:23. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:24.

[125]. In one aspect, one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 19 to 24 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEO ID NO:3. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:50, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:54 or SEQ ID NO:55. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1973 to 2006 of SEQ ID NO:3. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEO ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tagclosing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length.

[126]. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:7, SEQ ID NO:31, SEQ ID NO:56 or SEQ ID NO:57. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:7. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEQ ID NO:31.

[127]. In one aspect, one of said at least two tagged amplification oligomers comprises a target

hybridizing region that is from 20 to 25 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4. In another aspect, target-hybridizing region of said tagged amplification oligomer may contain one or more of a nucleotide addition, a nucleotide deletion, a nucleotide mismatch, a modified nucleotide or a combination thereof relative to a corresponding residue of said target nucleic acid. In another aspect, said target hybridizing region comprises at least 95% sequence identity to SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 or SEQ ID NO:62. In another aspect, said target hybridizing region is 15 to 30 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to a region of a target nucleic acid corresponding to residues 1994 to 2017 of SEQ ID NO:4. In one aspect, said tag region of said tagged amplification oligomer is from 10 to 25 nucleobases in length. In another aspect, said tag region is from 18 to 24 nucleobases in length. In another aspect, said tag region is 19 nucleobases in length. In another aspect, said tag region contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a sequence that is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said tagged amplification oligomer further comprises a tag-closing region, said tag-closing region comprising a nucleotide sequence that is substantially complementary to a portion of said target hybridizing region of said tagged-amplification oligomer. In another aspect, said tag-closing region is 3 to 20 nucleobases in length. In one aspect, said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:8, or SEQ ID NO:63. In another aspect, said tagged amplification oligomer comprises 100% sequence identity to SEO ID NO:8.

[128]. In one aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:5 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:7 and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:6 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that

is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer that is 100% identical to SEQ ID NO:7 and a tagged amplification oligomer that is 100% identical to SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:7 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer that is 100% identical to SEQ ID NO:6 and a tagged amplification oligomer that is 100% identical to SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:8 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer that is 100% identical to SEQ ID NO:6 and a tagged amplification oligomer that is 100% identical to SEQ ID NO:7. In another aspect, said at least two tagged amplification oligomers comprises four tagged amplification oligomers selected from the group consisting of: a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer that is at least 95% identical to SEO ID NO:6, a tagged amplification oligomer that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer that is 100% identical to SEQ ID NO:7, a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, and a tagged amplification oligomer that is 100% identical to SEO ID NO:8.

[129]. In one aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:5 and a second tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:6 and a second tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:6, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:6, SEQ ID NO:6, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:6 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises SEQ ID NO:8 and a second tagged amplification oligomer that is at least 95% identical to SEQ ID NO:8, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:9, SEQ ID NO:6, or SEQ ID NO:7, or that is 100% identical to SEQ ID NO:9.

NO:5, SEQ ID NO:6 or SEQ ID NO:7. In another aspect, said at least two tagged amplification oligomers comprises three tagged amplification oligomers each of the three individually being a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8. In another aspect, said at least two tagged amplification oligomers comprises four tagged amplification oligomers each of the four individually being a tagged amplification oligomer that is at least 95% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:6, SEQ ID NO:7, or SEQ ID NO:8, or that is 100% identical to SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8.

[130]. In one embodiment, said composition further comprises at least one blocker oligomer. In one aspect, one of said at least one blocker oligomer is at least 95% identical to SEQ ID NO:9 or SEQ ID NO:10. In another aspect, one of said at least one blocker oligomer is at least 95% identical to SEQ ID NO:9 or SEQ ID NO:10. In another aspect, said at least one blocker oligomer is two blocker oligomers, one of which is at least 95% identical to SEQ ID NO:9; or one of which is at least 95% identical to SEQ ID NO:9; or one of which is at least 95% identical to SEQ ID NO:9 and another of which is at least 95% identical to SEQ ID NO:9; or one of which is 100% identical to SEQ ID NO:10; or one of which is at least 95% identical to SEQ ID NO:9 and another of which is 100% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is at least 95% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is at least 95% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is 100% identical to SEQ ID NO:10; or one of which is 100% identical to SEQ ID NO:9 and another of which is 100% identical to SEQ ID NO:10.

[131]. In one aspect, said method at step (b.) results in at least one initial amplification product containing a tag sequence, or complement thereof. In another aspect, there is provided at least initial amplification product with a nucleotide sequence comprising a tag sequence region, wherein said tag sequence is from 10 to 25 nucleobases in length. In another aspect said initial amplification product comprises a tag sequence that is from 18 to 24 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that is 19 nucleobases in length. In another aspect, said initial amplification product comprises a tag sequence that contains a sequence that is substantially complementary to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33, or is substantially identical to one of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said initial amplification product comprises a tag sequence that contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof. In one aspect, said initial amplification product has a 3' end defined by a blocker oligomer. In another aspect, said initial

amplification product has a 3' end that is not defined by a blocker oligomer.

[132]. In one embodiment of the current invention, there is provided a method for the in vitro amplification of an initial amplification product, comprising the steps of (c.) contacting the sample with at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at the same time point. In one aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to the sample and a single amplification reaction is performed. In this aspect, steps (b.) and (d.) are combined to provide suitable conditions for performing an amplification reaction to generate an initial amplification product and for performing an amplification reaction generate a second amplification product therefrom. Though all of the amplification oligomers are present in this combined amplification reaction aspect, the tag-targeting amplification oligomers are configured to not participate in amplification until a tagged amplification oligomer is incorporated into an initial amplification product. In another aspect, the oligomers of step (c.) and the oligomers of step (a.) are contacted to said sample at separate time points. In one aspect, the method comprises the step of (c.) providing to an amplification product generated in step (b.) a second reaction mixture comprising a tag-targeting amplification oligomer consisting essentially of a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said tagged amplification oligomer, and comprising a second amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction. In one aspect for generating an amplification product, said sample is contacted with a first additional amplification oligomer that is a tagtargeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag region of said, or complement thereof in an initial amplification oligomer, and is contacted with a second additional amplification oligomer. In another aspect, said tag-targeting amplification oligomer comprises a target hybridization region that is 18 to 24 nucleobases in length. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:13. In another aspect, said tag-targeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:16. In another aspect, said tagtargeting amplification oligomer comprises a sequence that is 100% identical to SEQ ID NO:33. In one aspect, said second amplification oligomer is a primer amplification oligomer, a promoter based amplification oligomer or a tagged amplification oligomer. In another aspect, said second amplification oligomer is a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to

50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 SEQ ID NO:1. In another aspect, said promoter-based amplification oligomer comprises a target hybridizing region that is at least 95% identical to SEQ ID NO:42or that is 100% identical to SEQ ID NO:42. In another aspect, said a promoter-based amplification is at least 95% identical to SEQ ID NO:15 or is 100% identical to SEQ ID NO:15.

- [133]. In one embodiment there is further provided an analysis step, wherein an initial amplification product, an amplification product or both are analyzed. In one aspect, said analysis comprises a detection step to identify the presence or absence of an initial amplification product from an amplification reaction in step (b.), an amplification product generated in step (d.) or both initial amplification and amplification products generated in steps (b.) and (d.). In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is at least 95% identical to SEQ ID NO:14. In another aspect, said detection step is a probe-based detection step. In another aspect, said probe-based detection step uses a probe with a nucleotide sequence that is 100% identical to SEQ ID NO:14.
- [134]. In one embodiment, there is provided compositions, reaction mixtures and methods comprising at least one target capture oligomer. In one aspect, the target capture oligomers are also blocker oligomers. Preferred target capture oligomers include, but are not limited to SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:28, SEQ ID NO:29, SEQ ID NO:64, SEQ ID NO:65 and combinations thereof.

DETAILED DESCRIPTION

- [135]. Disclosed are compositions, reaction mixtures, kits and methods for amplifying *Mollicutes* nucleic acids from a sample; specifically sequences of a 23S rRNA or genes encoding a 23S rRNA. Amplified nucleic acids are then useful for a subsequent analysis. The compositions, reaction mixtures, kits and methods provide oligonucleotide sequences that recognize target sequences of 23S rRNA or their complementary sequences, or genes encoding 23S rRNA or their complementary sequences from a variety of species of *Mollicutes*. Such oligonucleotides include tagged amplification oligomers, blocker oligomers, tag-targeting amplification oligomers and promoter-based amplification oligomers; and in some embodiments further include target capture oligomers and detection probe oligomers. Reaction mixtures include target capture reagents, lysis reagents and amplification reagents. All or some of the oligomer compositions may be present in one or more of such reaction mixtures.
- [136]. The methods include performing a nucleic acid amplification of *Mollicutes* sequences. Amplified nucleic acids are then useful for a variety of subsequent analysis methods, including, but not limited to

nucleic acid detection, for example by specifically hybridizing the amplified product with a nucleic acid probe that provides a signal to indicate the presence of a *Mollicutes* in the sample. The amplification step includes contacting the sample with one or more amplification oligomers specific for a target sequence in 23S rRNA to produce an amplified product if a *Mollicutes* nucleic acid is present in the sample. Amplification synthesizes additional copies of the target sequence or its complement by using at least one nucleic acid polymerase to extend the sequence from an amplification oligomer (a primer) using a template strand. One embodiment for detecting the amplified product uses a hybridizing step that includes contacting the amplified product with at least one probe specific for a sequence amplified by the selected amplification oligomers, e.g., a sequence contained in the target sequence flanked by a pair of selected primers.

[137]. "Amplification" refers to any known procedure for obtaining multiple copies of a target nucleic acid sequence or its complement or fragments thereof. The multiple copies may be referred to as amplicons or amplification products. Amplification of "fragments" refers to production of an amplified nucleic acid that contains less than the complete target nucleic acid or its complement, e.g., produced by using an amplification oligonucleotide that hybridizes to, and initiates polymerization from, an internal position of the target nucleic acid. Known amplification methods include, for example, replicase-mediated amplification, polymerase chain reaction (PCR), ligase chain reaction (LCR), strand-displacement amplification (SDA), and transcription-mediated or transcription-associated amplification. Replicasemediated amplification uses self-replicating RNA molecules, and a replicase such as OB-replicase (e.g., US Pat. No. 4,786,600). PCR amplification uses a DNA polymerase, pairs of primers, and thermal cycling to synthesize multiple copies of two complementary strands of dsDNA or from a cDNA (e.g., US Pat. Nos. 4,683,195, 4,683,202, and 4,800,159). LCR amplification uses four or more different oligonucleotides to amplify a target and its complementary strand by using multiple cycles of hybridization, ligation, and denaturation (e.g., US Pat. No. 5,427,930 and US Pat. No. 5,516,663). SDA uses a primer that contains a recognition site for a restriction endonuclease and an endonuclease that nicks one strand of a hemimodified DNA duplex that includes the target sequence, whereby amplification occurs in a series of primer extension and strand displacement steps (e.g., US Pat. No. 5,422,252; US Pat. No. 5,547,861; and US 5,648,211). Those skilled in the art will know how to provide suitable conditions for performing these amplification reactions.

[138]. In the examples below, the nucleic acid of one or more species of the class *Mollicutes* is amplified by a transcription-based amplification technique. One transcription-based amplification system is transcription-mediated amplification (TMA), which employs an RNA polymerase to produce multiple RNA transcripts of a target region. Exemplary TMA amplification methods are described in U.S. Pat. Nos.

4,868,105; 5,124,246; 5,130,238; 5,399,491; 5,437,990; 5,480,784; 5,554,516; and 7,374,885; and PCT Pub. Nos. WO 88/01302; WO 88/10315 and WO 95/03430. The TMA reaction used below is a single primer TMA reaction as is described in U.S. Pat. No. 7,374,885. In general, the single-primer TMA methods use a primer oligomer, a modified promoter-based oligomer (or "promoter-provider oligomer") that is modified to prevent the initiation of DNA synthesis from its 3' end (e.g., by including a 3'-blocking moiety) and, optionally, a blocker oligomer to terminate elongation of a cDNA from the target strand. Promoter-based oligomers provide an oligonucleotide sequence that is recognized by an RNA polymerase. This single primer TMA method synthesizes multiple copies of a target sequence and includes the steps of treating a target RNA that contains a target sequence with a priming oligomer and a binding molecule, where the primer hybridizes to the 3' end of the target strand. RT initiates primer extension from the 3' end of the primer to produce a cDNA which is in a duplex with the target strand (e.g., RNA:cDNA). When a blocker oligomer, is used in the reaction, it binds to the target nucleic acid adjacent near the user designated 5' end of the target sequence. When the primer is extended by DNA polymerase activity of RT to produce cDNA, the 3' end of the cDNA is determined by the position of the blocker oligomer because polymerization stops when the primer extension product reaches the binding molecule bound to the target strand. Thus, the 3' end of the cDNA is complementary to the 5' end of the target sequence. The RNA:cDNA duplex is separated when RNase (e.g., RNase H of RT) degrades the RNA strand, although those skilled in the art will appreciate that any form of strand separation may be used. Then, the promoterprovider oligomer hybridizes to the cDNA near the 3' end of the cDNA strand. The promoter-provider oligomer includes a 5' promoter sequence for an RNA polymerase and a 3' target hybridizing region complementary to a sequence in the 3' region of the cDNA. The promoter-provider oligomer also has a modified 3' end that includes a blocking moiety that prevents initiation of DNA synthesis from the 3' end of the promoter-provider oligomer. In the promoter-provider:cDNA duplex, the 3'-end of the cDNA is extended by DNA polymerase activity of RT using the promoter oligomer as a template to add a promoter sequence to the cDNA and create a functional double-stranded promoter. An RNA polymerase specific for the promoter sequence then binds to the functional promoter and transcribes multiple RNA transcripts complementary to the cDNA and substantially identical to the target region sequence that was amplified from the initial target strand. The resulting amplified RNA can then cycle through the process again by binding the primer and serving as a template for further cDNA production, ultimately producing many amplicons from the initial target nucleic acid present in the sample. Some embodiments of the singleprimer transcription-associated amplification method do not include the blocking oligomer and, therefore, the cDNA product made from the primer has an indeterminate 3' end, but the amplification steps proceed substantially as described above for all other steps.

[139]. The current invention provides compositions and methods for uniplex and multiplex amplification

reactions. Assays for the nucleic acid amplification of *Mollicutes* nucleic acid in a sample source oftentimes show false positive or false negative results caused by contaminating from the environment and non-specific binding. Partly, these problems arise because of contaminating organisms present in the assay reagents or carried by the operators. For example, when the sample source is a bioreactor and the assay is to amplify *Mollicutes* common to such a source, then *M. orale M. salivarium M. pneumoniae* carried in by the assay operator can contaminate the assay. Partly too, laboratory environments where assays are performed and laboratory reagents can carry contaminating organisms that will non-specifically bind the assay reagents, leading to inefficient amplification of the target or amplification of non-target. False positives and false negatives are thus a significant challenge. Further challenging these assays when performed in multiplex are the deficiencies common to multiplex reactions, e.g., spurious product formation, primer dimers, unbalanced amplification caused by more efficient amplifications outperforming the less efficient reactions and thereby suing the reagents. Herein, the amplification oligomers comprise at least one tag region to address these and other problems.

[140]. Below describes an adaptation of a reverse transcription-mediated amplification (rTMA), various aspects of which are disclosed in U.S. Pat. Appln. Pub. No. US 2006-0046265 A1. The reaction of this illustrative embodiment is initiated by treating an RNA target sequence in a nucleic acid sample with both a tagged amplification oligomer and, optionally a blocking oligomer. The tagged amplification oligomer includes a target hybridizing region that hybridizes to a 3'-end of the target sequence and a tag region situated 5' to the target hybridizing region. The blocking oligomer hybridizes to a target nucleic acid containing the target sequence in the vicinity of the 5'-end of the target sequence. Thus, the target nucleic acid forms a stable complex with the tagged amplification oligomer at the 3'-end of the target sequence and the terminating oligonucleotide located adjacent to or near the determined 5'-end of the target sequence prior to initiating a primer extension reaction. Unhybridized tagged amplification oligomers are then made unavailable for hybridization to a target sequence prior to initiating a primer extension reaction with the tagged priming oligonucleotide, preferably by inactivating and/or removing the unhybridized tagged priming oligonucleotide from the nucleic acid sample. Unhybridized tagged amplification oligomer that has been inactivated or removed from the system is then unavailable for unwanted hybridization to contaminating nucleic acids. In one example of removing unhybridized tagged amplification oligomer from a reaction mixture, the tagged amplification oligomer is hybridized to the target nucleic acid, and the tagged amplification oligomer:target nucleic acid complex is removed from the unhybridized tagged amplification oligomer using a wash step. In this example, the tagged amplification oligomer:target nucleic acid complex may be further complexed to a target capture oligomer and a solid support. In one example of inactivating the unhybridized tagged amplification oligomer, the tagged amplification oligomers further comprise a target-closing region. In this example, the target hybridizing region of the

tagged amplification oligomer hybridizes to target nucleic acid under a first set of conditions (e.g., stringency). Following the formation of the tagged amplification oligomer:target nucleic acid complex the unhybridized tagged amplification oligomer is inactivated under a second set of the conditions, thereby hybridizing the target closing region to the target hybridizing region of the unhybridized tagged amplification oligomer. The inactivated tagged amplification oligomer is then unavailable for hybridizing contaminating nucleic acids. A wash step may also be included to remove the inactivated tagged amplification oligomers from the assay.

[141]. An extension reaction is then initiated from the 3'-end of the tagged amplification oligomer with a DNA polymerase, e.g., reverse transcriptase, to produce an initial amplification product that includes the tag sequence. The initial amplification product is then separated from the target sequence using an enzyme that selectively degrades the target sequence (e.g., RNAse H activity). Next, the initial amplification product is treated with a promoter-based oligomer having a target hybridizing region and an RNA polymerase promoter region situated 5' to the target hybridizing region, thereby forming a promoter-based oligomer:initial amplification product hybrid. In the examples below the promoter-based oligomer is modified to prevent the initiation of DNA synthesis, preferably by situating a blocking moiety at the 3'-end of the promoter-based oligomer (e.g., nucleotide sequence having a 3'-to-5' orientation). The 3'-end of the initial amplification product is then extended to add a sequence complementary to the promoter, resulting in the formation of a double-stranded promoter sequence. Multiple copies of a RNA product complementary to at least a portion of the initial amplification product are then transcribed using an RNA polymerase, which recognizes the double-stranded promoter and initiates transcription therefrom. As a result, the nucleotide sequence of the RNA product is substantially identical to the nucleotide sequence of the target nucleic acid and to the complement of the nucleotide sequence of the target nucleic acid and to the complement of the nucleotide sequence of the target nucleic acid and to the complement of the nucleotide sequence of

[142]. The RNA products are then treated with a tag-targeting oligomer, which hybridizes to the complement of the tag sequence to form a tag-targeting oligomer: RNA product hybrid, and the 3'-end of the tag-targeting oligomer is extended with the DNA polymerase to produce an amplification product complementary to the RNA product. The DNA strand of this amplification product is then separated from the RNA strand of this amplification product using an enzyme that selectively degrades the first RNA product (e.g., RNAse H activity). The DNA strand of the amplification product is treated with the promoter-based oligomer, which hybridizes to the 3'-end of the second DNA primer extension product to form a promoter-based oligomer:amplification product hybrid. The promoter-based oligomer:amplification product hybrid then re-enters the amplification cycle, where transcription is initiated from the double-stranded promoter and the cycle continues, thereby providing amplification product of the target sequence.

[143]. Amplification product can then be used in a subsequent assay. One subsequent assay includes nucleic acid detection, preferably nucleic acid probe-based nucleic acid detection. The detection step may be performed using any of a variety of known ways to detect a signal specifically associated with the amplified target sequence, such as by hybridizing the amplification product with a labeled probe and detecting a signal resulting from the labeled probe. The detection step may also provide additional information on the amplified sequence, such as all or a portion of its nucleic acid base sequence. Detection may be performed after the amplification reaction is completed, or may be performed simultaneous with amplifying the target region, e.g., in real time. In one embodiment, the detection step allows homogeneous detection, e.g., detection of the hybridized probe without removal of unhybridized probe from the mixture (e.g., US Pat. Nos. 5,639,604 and 5,283,174). In embodiments that detect the amplified product near or at the end of the amplification step, a linear probe may be used to provide a signal to indicate hybridization of the probe to the amplified product. One example of such detection uses a luminescentally labeled probe that hybridizes to target nucleic acid. Luminescent label is then hydrolyzed from non-hybridized probe. Detection is performed by chemiluminescence using a luminometer. (e.g., WO 89/002476). In other embodiments that use real-time detection, the probe may be a hairpin probe, such as a molecular beacon, molecular torch, or hybridization switch probe, that is labeled with a reporter moiety that is detected when the probe binds to amplified product. Such probes may comprise target binding sequences and non-target binding sequences. Various forms of such probes have been described previously (e.g., US Pat. Nos. 5,118,801; 5,312,728; 5,925,517; 6,150,097; 6,849,412; 6,835,542; 6,534,274; and 6,361,945; and US Pub. Nos. 20060068417A1; and US Pub. No. 20060194240A1).

- [144]. To aid in understanding aspects of the invention, some terms used herein are described in more detail. All other scientific and technical terms used herein have the same meaning as commonly understood by those skilled in the relevant art, such as may be provided in *Dictionary of Microbiology and Molecular Biology*, 2nd ed. (Singleton et al., 1994, John Wiley & Sons, New York, NY), *The Harper Collins Dictionary of Biology* (Hale & Marham, 1991, Harper Perennial, New York, NY), and other references cited herein. Unless mentioned otherwise, the techniques employed or contemplated herein are standard methods well known to a person of ordinary skill in the art of molecular biology.
- [145]. It is to be noted that the term "a" or "an" entity refers to one or more of that entity; for example, "a nucleic acid," is understood to represent one or more nucleic acids. As such, the terms "a" (or "an"), "one or more," and "at least one" can be used interchangeably herein.
- [146]. "Sample" includes those that may contain one or more species from the class *Mollicutes* or components thereof, such as nucleic acids or fragments of nucleic acids wherein the source of that sample may contain an undesired species of *Mollicutes*. Samples include bioreactors; mammalian, avian, insect,

fish or plant cell lines; laboratory workspaces, laboratory equipment, cell-culture wares, laboratory reagents, biological manufacturing wares, pharmaceutical manufacturing wares, biological drug product, biological bulk drug substance, pharmaceutical drug product, pharmaceutical bulk drug substance and biological samples.

- [147]. "Nucleic acid" refers to a multimeric compound comprising two or more covalently bonded nucleosides or nucleoside analogs having nitrogenous heterocyclic bases, or base analogs, where the nucleosides are linked together by phosphodiester bonds or other linkages to form a polynucleotide. A "nucleotide" as used herein is a subunit of a nucleic acid consisting of a phosphate group, a 5-carbon sugar and a nitrogenous base. The 5-carbon sugar found in RNA is ribose. In DNA, the 5-carbon sugar is 2'-deoxyribose. The term also includes analogs of such subunits, such as a methoxy group at the 2' position of the ribose (2'-O-Me). As used herein, methoxy oligonucleotides containing "T" residues have a methoxy group at the 2' position of the ribose moiety, and a uracil at the base position of the nucleotide. A "non-nucleotide unit" as used herein is a unit that does not significantly participate in hybridization of a polymer. Such units must not, for example, participate in any significant hydrogen bonding with a nucleotide, and would exclude units having as a component one of the five nucleotide bases or analogs thereof.
- [148]. The term "polynucleotide" as used herein denotes a nucleic acid chain. Throughout this application, nucleic acids are designated by the 5'-terminus to the 3'-terminus. Standard nucleic acids, e.g., DNA and RNA, are typically synthesized "3'-to-5'," i.e., by the addition of nucleotides to the 5'-terminus of a growing nucleic acid.
- [149]. A "target nucleic acid" as used herein is a nucleic acid comprising a "target sequence" to be amplified. Target nucleic acids may be DNA or RNA as described herein, and may be either single-stranded or double-stranded. The target nucleic acid may include other sequences besides the target sequence, which may not be amplified. The preferred target nucleic acid herein is a nucleic acid from one or more species of the class *Mollicutes*.
- [150]. By "isolated" it is meant that a sample containing a target nucleic acid is taken from its natural milieu, but the term does not connote any degree of purification. "Separating" or "purifying" means that one or more components of a sample are removed or separated from other sample components. Sample components include target nucleic acids usually in a generally aqueous solution phase, which may also include cellular fragments, proteins, carbohydrates, lipids, and other nucleic acids. Separating or purifying removes at least 70%, of the target nucleic acid from other sample components.

[151]. The term "target sequence" as used herein refers to the particular nucleotide sequence of the target nucleic acid that is to be amplified and/or detected. The "target sequence" includes the sequences to which oligomers hybridize (e.g., tagged amplification oligomer, tag-target oligomers, promoter-based amplification oligomers, blocking oligomers, target-capture oligomers, detection oligomers). Where the target nucleic acid is originally single-stranded, the term "target sequence" will also refer to the sequence complementary to the "target sequence" as present in the target nucleic acid. Where the target nucleic acid is originally double-stranded, the term "target sequence" refers to both the sense (+) and antisense (-) strands. In choosing a target sequence, the skilled artisan will understand that a "unique" sequence should be chosen so as to distinguish between unrelated or closely related target nucleic acids.

- [152]. "Target binding region" is used herein to refer to the portion of an oligomer that is configured to hybridize with a target nucleic acid sequence. Preferably, the target binding regions are configured to specifically hybridize with a target nucleic acid sequence. Target binding regions may be 100% complementary to the portion of the target sequence to which they are hybridize; but not necessarily. Target-binding regions may also include inserted, deleted and/or substituted nucleotide residues relative to a target sequence. Less than 100% complementarity of a target binding region to a target sequence may arise, for example, when the target nucleic acid is a plurality of strains within a species or within a class, such as would be the case for an oligomer that hybridizes to a variety of the strains of *Mollicutes* class. It is understood that other reasons exist for configuring a target binding region to have less than 100% complementarity to a target nucleic acid. Target hybridizing regions are preferably about 10 to 50 nucleotides in length. In the current description, for some of the oligomers, the target hybridizing regions are described as being configured to hybridize to a region of a reference sequence. This type of description is not a limitation that an oligomer comprising such a target hybridizing region is one that is limited to hybridizing only to that reference sequence. Similarly, this description does not exclude target hybridizing regions that hybridize to the complementary strand of that illustrated by the reference sequence.
- [153]. The term "targets a sequence" as used herein in reference to a region of a *Mollicutes* nucleic acid refers to a process whereby an oligonucleotide hybridizes to the target sequence in a manner that allows for amplification as described herein. Preferably, the oligomer specifically hybridizes to the target sequence or group of target sequences.
- [154]. The term "fragment" as used herein in reference to the *Mollicutes* targeted nucleic acid sequence refers to a piece of contiguous nucleic acid. In certain embodiments, the fragment includes contiguous nucleotides from a *Mollicutes* 23S ribosomal RNA, wherein the number of 23S contiguous nucleotides in the fragment are less than that for the entire 23S.

[155]. The term "region" as used herein refers to a portion of a nucleic acid wherein said portion is smaller than the entire nucleic acid. In one example the nucleic acid in reference is an amplification oligomer, such as a tagged-amplification oligomer. Here, the term "region" may be used refer to the smaller target hybridizing-portion of the entire oligomer or the tag portion of the entire oligomer. Similarly, and also as example only, when the nucleic acid is a 23S ribosomal RNA, the term "region" may be used to refer to a smaller area of the nucleic acid, wherein the smaller area is targeted by one or more oligonucleotides of the invention. As a further non-limiting example, when the nucleic acid in reference is an initial amplification product, the term region may be used to refer to the smaller tag nucleotide sequence or complementary tag nucleotide sequence identified for hybridization by the target binding region of a tagged-amplification oligomer.

- [156]. The interchangeable terms "oligomer," "oligo" and "oligonucleotide" refer to a nucleic acid having generally less than 1,000 nucleotide (nt) residues, including polymers in a range having a lower limit of about 5 nt residues and an upper limit of about 500 to 900 nt residues. In some embodiments, oligonucleotides are in a size range having a lower limit of about 10 to 15 nt and an upper limit of about 50 to 600 nt, and other embodiments are in a range having a lower limit of about 15 to 20 nt and an upper limit of about 22 to 100 nt. These ranges are merely exemplary. Oligonucleotides may be purified from naturally occurring sources, or may be synthesized using any of a variety of well known enzymatic or chemical methods. The term oligonucleotide does not denote any particular function to the reagent; rather, it is used generically to cover all such reagents described herein. An oligonucleotide may serve various different functions. Herein, the oligomers referred to are primarily amplification oligomers such as primer oligomers, promoter-based amplification oligomers, tagged amplification oligomers, tag-targeting oligomers; target capture oligomers, detection oligomers and blocker oligomers. This list of oligomers is non-limiting, as is understood by the skilled artisan.
- [157]. As used herein, an oligonucleotide having a nucleic acid sequence "comprising" or "consisting of" or "consisting essentially of" a sequence selected from a group of specific sequences means that the oligonucleotide, as a basic and novel characteristic, is capable of stably hybridizing to a nucleic acid having the exact complement of one of the listed nucleic acid sequences of the group under stringent hybridization conditions. An exact complement includes the corresponding DNA or RNA sequence.
- [158]. "Consisting essentially of" is used to mean that additional component(s), composition(s) or method step(s) that do not materially change the basic and novel characteristics of the present invention may be included in the compositions or kits or methods of the present invention. Such characteristics include the ability to detect *Mollicutes* nucleic acid in a sample. Other characteristics include limited cross-reactivity with other Bacteria or mammalian nucleic acid and targeting 23S rRNA. Any

component(s), composition(s), or method step(s) that have a material effect on the basic and novel characteristics of the present invention would fall outside of this term.

[159]. As used herein, an oligonucleotide "substantially corresponding to" a specified nucleic acid sequence means that the referred to oligonucleotide is sufficiently similar to the reference nucleic acid sequence such that the oligonucleotide has similar hybridization properties to the reference nucleic acid sequence in that it would hybridize with the same target nucleic acid sequence under stringent hybridization conditions. One skilled in the art will understand that "substantially corresponding oligonucleotides" can vary from the referred to sequence and still hybridize to the same target nucleic acid sequence. It is also understood that a first nucleic acid corresponding to a second nucleic acid includes the complements thereof and includes the RNA and DNA thereof. This variation from the nucleic acid may be stated in terms of a percentage of identical bases within the sequence or the percentage of perfectly complementary bases between the probe or primer and its target sequence. Thus, an oligonucleotide "substantially corresponds" to a reference nucleic acid sequence if the nucleotide sequence of the first nucleic acid is from 100% to about 70% identical, or in some instances complementary, to the reference nucleic acid sequence. This range is inclusive of whole and partial numbers, e.g., 70%, 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, 79%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100%. In preferred embodiments, the percentage is from 100% to about 85%. In more preferred embodiments, this percentage can be from 100% to about 90%; in other preferred embodiments, this percentage is from 100% to about 95%. Similarly, a region of a nucleic acid or amplified nucleic acid can be referred to herein as corresponding to a reference nucleic acid sequence. One skilled in the art will understand the various modifications to the hybridization conditions that might be required at various percentages of complementarity to allow hybridization to a specific target sequence without causing an unacceptable level of non-specific hybridization.

[160]. As used herein, a "blocking moiety" is a substance used to "block" the 3'-terminus of an oligonucleotide or other nucleic acid so that it cannot be efficiently extended by a nucleic acid polymerase. Oligomers not intended for extension by a nucleic acid polymerase may include a blocker group that replaces the 3'OH to prevent enzyme-mediated extension of the oligomer in an amplification reaction. For example, blocked amplification oligomers and/or detection probes present during amplification may not have functional 3'OH and instead include one or more blocking groups located at or near the 3' end. In some embodiments a blocking group near the 3' end and may be within five residues of the 3' end and is sufficiently large to limit binding of a polymerase to the oligomer. In other embodiments a blocking group is covalently attached to the 3' terminus. Many different chemical groups may be used to block the 3' end, e.g., alkyl groups, non-nucleotide linkers, alkane-diol dideoxynucleotide residues, and cordycepin.

[161]. An "amplification oligomer" is an oligomer, at least the 3'-end of which is complementary to a target nucleic acid, and which hybridizes to a target nucleic acid, or its complement, and participates in a nucleic acid amplification reaction. An example of an amplification oligomer is a "primer" that hybridizes to a target nucleic acid and contains a 3' OH end that is extended by a polymerase in an amplification process. Another example of an amplification oligomer is an oligomer that is not extended by a polymerase (e.g., because it has a 3' blocked end) but participates in or facilitates amplification. For example, the 5' region of an amplification oligonucleotide may include a promoter sequence that is non-complementary to the target nucleic acid (which may be referred to as a "promoter-primer" or "promoter provider"). Those skilled in the art will understand that an amplification oligomer that functions as a primer may be modified to include a 5' promoter sequence, and thus function as a promoter-primer. Further an amplification oligomer that functions as a primer may be modified to include a 5' tag region, thereby incorporating into an initial amplification product or amplification product a universal sequence. Size ranges for the target hybridizing regions of amplification oligonucleotides include those that are about 10 to about 70 nt in length and are substantially complementary to a region of the target nucleic acid sequence (or a complementary strand thereof).

- [162]. As used herein, a "promoter" is a specific nucleic acid sequence that is recognized by a DNA-dependent RNA polymerase ("transcriptase") as a signal to bind to the nucleic acid and begin the transcription of RNA at a specific site.
- [163]. As used herein, a "promoter-provider" or "provider" refers to an oligonucleotide comprising first and second regions, and which is modified to prevent the initiation of DNA synthesis from its 3'-terminus. The "first region" or "target hybridizing region" of a promoter–provider oligonucleotide comprises a base sequence which hybridizes to a DNA template, where the hybridizing sequence is situated 3', but not necessarily adjacent to, a promoter region. The target hybridizing region of a promoter-based oligonucleotide is typically at least 10 nucleotides in length, and may extend up to 50 or more nucleotides in length. The "second region" or "promoter region" comprises a promoter sequence for an RNA polymerase. A promoter oligonucleotide is engineered so that it is incapable of being extended by an RNA- or DNA-dependent DNA polymerase, *e.g.*, reverse transcriptase, preferably comprising a blocking moiety at its 3'-terminus as described above. As referred to herein, a "T7 Provider" is a blocked promoter-provider oligonucleotide that provides an oligonucleotide sequence that is recognized by T7 RNA polymerase. A similar oligomer that lacks modification to the 3'-terminus is called a "promoter primer."
- [164]. As used herein, a "blocker oligonucleotide" is an oligonucleotide comprising a base sequence that is complementary to a region of the target nucleic acid in the vicinity of the 5'-end of the target sequence,

so as to "terminate" primer extension of a nascent nucleic acid that includes a priming oligonucleotide, thereby providing a defined 3'-end for the nascent nucleic acid strand.

- [165]. The term "initial amplification product" is used herein in reference to synthesis of a complementary strand of nucleic acid, wherein said synthesis incorporates into the newly synthesized strand a tag region sequence. In subsequent amplifications wherein the template nucleic acid comprises such a tag region sequence or complement thereof or the term "amplification product" is used.

 Referencing the TMA amplification described above, the initial amplification product is the product generated from the RNA template using the tagged amplification oligomer to incorporate the tag region into the initial amplification product. The amplification product is, therefore, the product that is generated following degradation of the original target nucleic acid from the new DNA strand, which for the above exemplary amplification would include product generated using the promoter-based amplification oligomer, product generated by the RNA polymerase and product generated using the tag-targeting amplification oligomers. This detailed description is used to help with understanding of these terms. The use of two terms for the nucleic acids synthesized in the amplification reaction is merely for convenience and does not limit the current invention. Similarly, ordinarily skilled artisans will understand how these terms apply with other isothermal and cyclical amplification methods.
- [166]. "Probe," "detection probe" or "detection oligonucleotide" are terms referring to a nucleic acid oligomer that hybridizes specifically to a target sequence in a nucleic acid, or in an amplified nucleic acid, under conditions that promote hybridization to allow detection of the target sequence or amplified nucleic acid. Detection may either be direct (e.g., a probe hybridized directly to its target sequence) or indirect (e.g., a probe linked to its target via an intermediate molecular structure). Probes may be DNA, RNA, analogs thereof or combinations thereof and they may be labeled or unlabeled. A probe's "target sequence" generally refers to a smaller nucleic acid sequence region within a larger nucleic acid sequence that hybridizes specifically to at least a portion of a probe oligomer by standard base pairing. A probe may comprise target-specific sequences and other sequences that contribute to the three-dimensional conformation of the probe (e.g., U.S. Pat. Nos. 5,118,801; 5,312,728; 6,849,412; 6,835,542; 6,534,274; and 6,361,945; and U.S. Pub. No. 20060068417).
- [167]. As used herein, a "target capture oligomer" or "target capture probe" refers to a nucleic acid oligomer that specifically hybridizes to a target sequence in a target nucleic acid by standard base pairing and joins to a binding partner on an immobilized probe to capture the target nucleic acid to a support. One example of a target capture oligomer includes two binding regions: the first being a target-hybridizing region and the second being an immobilized probe-binding region. The two regions may be present on two different oligomers joined together by one or more linkers, or the two regions may be on the same

oligomer. Another embodiment of a capture oligomer uses a target binding region that includes random or non-random poly-GU, poly-GT, or poly U sequences to bind non-specifically to a target nucleic acid and link it to an immobilized probe on a support. The immobilized probe-binding region may be a heteropolymeric nucleotide sequence or a homopolymeric nucleotide sequence. Exemplary heteropolymeric sequences include dT.sub.3/dT.sub.30 nucleotide sequences. Exemplary homopolymeric nucleotide sequences include dT.sub.30, dA.sub.30, dC.sub.30 or dG.sub.30 nucleotide sequences. (See e.g., U.S. Pat. No. 6,110,678, 6,280,952, and 6,534,273 and U.S. Pub. No. 2008-0286775 A1).

[168]. As used herein, an "immobilized oligonucleotide", "immobilized probe" or "immobilized nucleic acid" refers to a nucleic acid binding partner that joins a capture oligomer to a support, directly or indirectly. An immobilized probe joined to a support facilitates separation of a capture probe bound target from unbound material in a sample. One embodiment of an immobilized probe is an oligomer joined to a support that facilitates separation of bound target sequence from unbound material in a sample. Supports may include known materials, such as matrices and particles free in solution, which may be made of nitrocellulose, nylon, glass, polyacrylate, mixed polymers, polystyrene, silane, polypropylene, metal, or other compositions, of which one embodiment is magnetically attractable particles. Supports may be monodisperse magnetic spheres (e.g., uniform size \pm 5%), to which an immobilized probe is joined directly (via covalent linkage, chelation, or ionic interaction), or indirectly (via one or more linkers), where the linkage or interaction between the probe and support is stable during hybridization conditions.

[169]. By "complementary" is meant that the nucleotide sequences of similar regions of two single-stranded nucleic acids, or to different regions of the same single-stranded nucleic acid have a nucleotide base composition that allow the single-stranded regions to hybridize together in a stable double-stranded hydrogen-bonded region under stringent hybridization or amplification conditions. Sequences that hybridize to each other may be completely complementary or partially complementary to the intended target sequence by standard nucleic acid base pairing (e.g. G:C, A:T or A:U pairing). By "sufficiently complementary" or "substantially complementary" is meant a contiguous sequence that is capable of hybridizing to another sequence by hydrogen bonding between a series of complementary bases, which may be complementary at each position in the sequence by standard base pairing or may contain one or more residues, including abasic residues, that are not complementary. Sufficiently complementary contiguous bases are at least 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100% complementary to the target sequence to which the target hybridizing region hybridizes. Skilled artisans know that such complementarity need not include only whole numbers (e.g., a 27 nucleotide oligomer having 22 nucleotides that are complementary to a target sequence will have an 81.48% complementarity to the target). Sequences that are "sufficiently

complementary" allow stable hybridization of a nucleic acid oligomer with its target sequence under appropriate hybridization conditions, even if the sequences are not completely complementary. The same applies to sequence identity, wherein one nucleic acid has a nucleic acid sequence that is all or partially identical to a nucleotide sequence within another nucleic acid molecule.

- [170]. By "preferentially hybridize" or "specifically hybridize" is meant that under stringent hybridization assay conditions, oligomers hybridize to the reference target sequence, or replicates thereof, to form stable oligomer:target hybrids, while at the same time formation of stable oligomer:non-target hybrids is minimized. Thus, an oligomer hybridizes to a target sequence or replicate thereof to a sufficiently greater extent than to a non-target sequence, to enable one having ordinary skill in the art to accurately quantitate the RNA replicates or complementary DNA (cDNA) of the target sequence formed during the amplification. Compositions that are configured to specifically hybridize to a reference sequence are not limited to hybridizing only that reference sequence (e.g., only the *Mycoplasma hominis* species exemplified by SEQ ID NO:1). Rather, all that is meant is that such compositions have a nucleotide sequence that specifically hybridizes to the reference sequence but the composition will hybridize other species sequences (e.g., oligomers configured to specifically hybridize to a region of SEQ ID NO:1 will also hybridize the corresponding region in other *Mycoplasma* species). Compositions that specifically hybridize to a region of a target nucleic acid may have one or more of a nucleotide addition, deletion, mismatch, or modified nucleotide relative to a corresponding residue on the target nucleic acid.
- [171]. "Sample preparation" refers to any steps or method that treats a sample for subsequent amplification and/or detection of *Mollicutes* nucleic acids present in the sample. Samples may be complex mixtures of components of which the target nucleic acid is a minority component. Sample preparation may include any known method of concentrating components, such as microbes or nucleic acids, from a larger sample volume, such as by filtration of airborne or waterborne particles from a larger volume sample or by isolation of microbes from a sample by using standard microbiology methods. Sample preparation may include physical disruption and/or chemical lysis of cellular components to release intracellular components into a substantially aqueous or organic phase and removal of debris, such as by using filtration, centrifugation or adsorption. Sample preparation may include use of one or more target capture oligomers.
- [172]. The term "specificity," in the context of an amplification and/or detection system, is used herein to refer to the characteristic of the system which describes its ability to distinguish between target and non-target sequences dependent on sequence and assay conditions. In terms of nucleic acid amplification, specificity generally refers to the ratio of the number of specific amplicons produced to the number of side-

products (e.g., the signal-to-noise ratio). In terms of detection, specificity generally refers to the ratio of signal produced from target nucleic acids to signal produced from non-target nucleic acids.

- [173]. The term "sensitivity" is used herein to refer to the precision with which a nucleic acid amplification reaction can be detected or quantitated. The sensitivity of an amplification reaction is generally a measure of the smallest copy number of the target nucleic acid that can be reliably detected in the amplification system, and will depend, for example, on the detection assay being employed, and the specificity of the amplification reaction, e.g., the ratio of specific amplicons to side-products.
- [174]. As used herein, the term "relative fluorescence unit" ("RFU") is an arbitrary unit of measurement of fluorescence intensity. RFU varies with the characteristics of the detection means used for the measurement.
- [175]. As used herein, the term "TTime" is the threshold time or time of emergence of signal in a real-time plot of the assay data. TTime values estimate the time at which a particular threshold indicating amplicon production is passed in a real-time amplification reaction. TTime and an algorithm for calculating and using TTime values are described in U.S. Pub. No. 2006/0276972, paragraphs [0517] through [0538].
- [176]. In a preferred embodiment, a 23S rRNA or a gene encoding the 23S rRNA of a species in the class Mollicutes is selectively separated from other sample components by specifically hybridizing the Mollicutes nucleic acid to a target capture oligomer to form a target sequence:target capture probe complex that is then separated from sample components. A preferred method of specific target capture binds the Mollicutes nucleic acid target sequence:target capture probe complex to an immobilized probe to form a nucleic acid target sequence:target capture probe:immobilized probe complex that is separated from the sample and, optionally, washed to remove non-target sample components. Also removed in this preferred embodiment is unhybridized tagged amplification oligomer, and more preferably inactivated unhybridized tagged amplification oligomer. The target capture probe preferably includes a a specific binding partner that attaches the capture probe with its bound target sequence to a solid support, to facilitate separating the target sequence from the sample components. In a preferred embodiment, the specific binding partner of the capture probe is a 3' heteropolymeric or homopolymeric tail sequence that is not complementary to the target sequence but that hybridizes to a complementary sequence on an immobilized probe attached to a solid support. Target capture preferably occurs in a solution phase mixture that contains one or more target capture oligomers that hybridize specifically to the target nucleic acid under hybridizing conditions. usually at a temperature higher than the Tm of the tail sequence:immobilized probe sequence duplex. Then, the target sequence:target capture probe complex is captured by adjusting the hybridization

conditions so that the capture probe tail hybridizes to the immobilized probe, and the entire complex on the solid support is then separated from other sample components. Preferred embodiments use a particulate solid support, such as paramagnetic beads, so that particles with the attached target sequence:target capture probe:immobilized probe complex may be suspended in a washing solution and retrieved from the washing solution, preferably by using magnetic attraction. In a more preferred embodiment, the target capture occurs in solution phase and further comprises one or more tagged amplification oligomers, which bind to the target nucleic acid, thus forming part of the captured oligomer complex bound to the solid support. In this preferred embodiment, the tagged amplification oligomer is maintained with its target during the removal step, while unhybridized and/or inactivated unhybridized tagged amplification oligomer is removed, preferably via a wash step.

[177]. The following examples illustrate some of the embodiments of the invention for amplification of Mollicutes 23S rRNA target sequences. A plurality of tagged amplification oligomers were prepared and tested for sensitivity and specificity in amplifying and detecting different species in the class Mollicutes without interference from common cross reactors. The target species tested in these following examples include: M. orale; M. arginini; M. hominis; M. synoviae; M. arthritidis; M. hyorhinis; A. laidlawii; Ureaplasma; S. citri; M. pneumoniae; M. gallisepticum; M. pirum; M. salivarium; and M. fermentans. The following species were used as challenge organisms: E. coli; C. perfringens; S. bovis; S. epidermidis; S. aureus; P. acnes; E. faecalis; B. subtillis; Clostridium sporogenes; Lactobacillus casei; Corynebacterium pseudodiphthericum; P. Aeruginosa; M. luteus; and L. pneumophila. A Chinese Hamster Ovary (CHO) cell line was also used as a challenge organism. These target and challenge organisms are available from a variety of well known sources, including Poultry Diagnostic Research Center (Athens, GA) and biological depositories such as American Type Cell Culture (ATCC, Manassas, VA) to name a few.

[178]. Example 1: Amplification of Target Nucleic Acid from Various Species in the class *Mollicutes*; Sensitivity and Specificity.

[179]. A first target capture reaction mixture was prepared to comprise a target capture oligomer (SEQ ID NO:29) and one of four tagged amplification oligomers (SEQ ID NO:73; SEQ ID NO:74; SEQ ID NO:75; or SEQ ID NO:76). The target capture oligomer was configured to hybridize to a target species nucleic acid at a position that also allows the target capture oligomer to perform as a blocker oligomer when these tagged amplification oligomers are used for generating an initial amplification product. The amplification reaction mixture comprised a promoter-based amplification oligomer (SEQ ID NO:15) and a tag-targeting amplification oligomer (SEQ ID NO:66). The target capture and amplification reactions were performed as is generally described herein. Briefly, lysates containing either 1000 copies of *M. fermentans* in the presence of either 1E+6 copies of the challenge organism *S. bovis* or 1E+6 copies of the challenge

organism *E. coli* were incubated in target capture reagent for 60 minutes at 42.deg.C followed by 15 minutes at room temp. A capture and was step was performed using magnetic beads and a KingFisher ml purification system. Following target capture, amplification reaction mixture was added to each of the reaction conditions and a TMA reaction was performed at 42.deg.C using a molecular torch (SEQ ID NO:14). Positive cut-off was set at 1000 RFU. Three of the four tagged amplification oligomers showed good specificity for *M. fermentans* target at 1000 copies in the presence of both challenge organisms; and SEQ ID NO:76 showed was the best performer in this example. The results for SEQ ID NO:73 showed cross reactivity with *S. bovis*, having an average TTime of about 20.71 minutes (±7.1).

- [180]. Further target capture and amplification reactions were prepared wherein the target capture mixtures comprised a target capture oligomer (SEQ ID NO:25) and one of the following tagged amplification oligomers (SEQ ID NOS:17, 18, 19, 20, 21, 22, 23 or 24) combined separately with each of the following blocker oligomers (SEQ ID NOS:25 or 26). The amplification reagent comprised a promoter-based amplification oligomer (SEQ ID NO:15) and a tag targeting oligomer (SEQ ID NO:16). Target nucleic acid was 0, 1E+3 or 1E+4 copies of *M. fermentans* nucleic acid. Target capture was performed as is generally described herein. Briefly, each of the samples were incubated at 60.deg.C for 15 minutes followed by room temperature for 5 minutes. Target capture reagent was added to the sample and these mixtures were mixed, incubated and washed. Following target capture, amplification reaction mix was added to each of the reaction conditions and a TMA reaction was performed at 42.deg.C in the presence of SEQ ID NO:27, which is a molecular torch. *M. fermentans* target nucleic acid was detected at 1E+3 and 1E+4 copies in each reaction condition, with TTimes ranging from about 12 to about 27 minutes. However, there were also false positives in samples comprising SEQ ID NOS:17, 18, 19 and 21 tagged amplification oligomers.
- [181]. Example 2: Amplification of Target Nucleic Acid from Various Species in the class *Mollicutes*; Sensitivity in Uniplex and Duplex Reactions.
- [182]. Additional amplification and detection conditions were prepared to test the tagged amplification oligomer SEQ ID NO:76 in a uniplex reaction and in a duplex reaction. Two uniplex target capture reaction mixtures were prepared, the first comprising SEQ ID NO:29 and SEQ ID NO:76; and the second comprising SEQ ID NO:29 and SEQ ID NO:67 as a tagged amplification oligomer. A duplex target capture reaction mixture was prepared to comprise SEQ ID NO:29, SEQ ID NO:76 and SEQ ID NO:67. The amplification reaction mixture comprised SEQ ID NOS:15 & 66. The presence or absence of an amplification product was determined using SEQ ID NO:14. Target was 1E+2 or 1E+3 copies of each of *M. fermentans* (SEQ ID NO:76 uniplex reaction) or *M. gallisepticum*; *M. pneumoniae* or *Ureaplasma* (SEQ ID NO:67 uniplex reaction) and 1E+2 or 1E+3 copies of each of *M. fermentans*, *M. hyorhinus* and

M. gallisepticum for the duplex reaction. Target capture and TMA were performed as is generally described herein. Positive samples were determined as those with RFUs above 1000. The uniplex reactions showed sensitivity to 1E+2 copies of all target nucleic acid tested. The duplex reaction also showed sensitivity to 1E+2 for M. fermentans and M. gallisepticum, but did not detect M. hyorhinus. Thus, the two tagged amplification oligomers did not interfere with each other and did not increase background, however, the oligomers did not detect one of the target species.

- [183]. Example 3: Amplification of Target Nucleic Acid from Various Species in the class *Mollicutes*; Sensitivity in Duplex Reactions.
- [184]. Additional tagged amplification oligomers were prepared and tested for sensitivity to *M. hyorhinus* sensitivity. *M gallisepticum* was used as positive control. Four target capture reaction mixtures were prepared to comprise two target capture oligomers (SEQ ID NOS:28 & 29); a first tagged amplification oligomer (SEQ ID NO:67) and one of four second tagged amplification oligomers (SEQ ID NO:68; SEQ ID NO:74; SEQ ID NO:75; or SEQ ID NO:76). Both target capture oligomers were configured to hybridize to a target species nucleic acid at a position that also allows the target capture oligomer to perform as a blocker oligomer. The amplification reaction mixture comprised a promoter-based amplification oligomer (SEQ ID NO:15) and a tag-targeting amplification oligomer (SEQ ID NO:66). The presence or absence of an amplification product was determined using SEQ ID NO:14, a molecular torch. Target was 1E+2 or 1E+3 copies of *M. hyorhinus*. Positive control was 1E+2 or 1E+3 copies of *M. gallisepticum*. Negative control was water. Target capture and TMA amplification was performed as is generally described herein. All duplex reactions showed sensitivity to *M. gallisepticum* down to 1E+2 copies per reaction. Likewise, all duplex reactions amplified 1E+3 copies of *M. hyorhinus*, though only the duplex reactions comprising SEQ ID NO:68; SEQ ID NO:74 or SEQ ID NO:75 could amplify *M. hyorhinus* to 1E+2 copies.
- [185]. Example 4: Amplification of Target Nucleic Acid from Various Species in the Class *Mollicutes*; Sensitivity in Multiplex Reactions.
- [186]. Target capture reagent mixtures were prepared comprising three tagged amplification oligomers (SEQ ID NOS:67, 68 & 69) and two target capture oligomers (SEQ ID NOS:28 & 29). Samples containing target nucleic acids were prepared as follows, 0, 1E+2 or 1E+3 copies of *M. hyorhinus*, *M. arginini*, *M. hominus*, *M. orales*, *M. gallisepticum*, *M. pneumoniae*, *M. fermentans* or *A. laidlawii*. Target capture was performed as is generally described herein. Amplification reagent comprised SEQ ID NO:15 and SEQ ID NO:66 as promoter-based amplification oligomer and tag targeting amplification oligomer, respectively. TMA amplification was also performed as is generally described herein and detection of

amplified product was performed using SEQ ID NO:14. In this multiplex reaction, all target nucleic acids were amplified well at 1E+3 copies of nucleic acid per reaction with an average TTime of about 22.51 minutes (±1.74). Similarly, all target nucleic acids at 1E+2 copies per reaction were amplified in these multiplex reactions average TTime of about 26.2 minutes (±1.78). There were no false positives.

[187]. In another multiplex reaction using the target capture reagent and the amplification reagent described directly above (target capture reagent comprises SEQ ID NOS:28, 29, 67, 68 & 69; and amplification reagent comprises SEQ ID NOS:15 & 66). In this reaction, the target nucleic acid was extracted from the following four organisms: *M. arthritidis*, *M. pirum*, *M. salvarium* and *M. gallisepticum*. Positive control was 1E+3 copies of 23S rRNA from *M. gallisepticum*. Samples were prepared at 1 CFU per reaction. The presence or absence of target nucleic acid was determined in real time using SEQ ID NO:14. In this multiplex reaction, all target nucleic acids were amplified at 1 CFU (approximately 1E+3 copies per CFU). The positive control was also positive and there were no false positives.

[188]. Example 5: Amplification of Target Nucleic Acid from Various Species in the Class *Mollicutes* in the Presence of Challenge Nucleic Acids; Sensitivity in Multiplex Reactions.

[189]. Target capture reagent mixtures were prepared comprising three tagged amplification oligomers (SEQ ID NOS:67, 68 & 69) and two target capture oligomers (SEQ ID NOS:28 & 29). Samples contained 1E+8 copies of nucleic acid from one of the following challenge organisms: S.epidermidis, S. bovis, M. luteus, B. subtillis, E. coli, L. pneumophilia, P. aeruginosa, Chinese Hamster Ovary (CHO) cell, Clostridium sporogenes, Lactobacillus casel, or Corynebacterium pseudodiphtheriticum. These samples were prepared in duplicate and one of the duplicates contained 1E+3 copies of M. gallisepticum and the other of the duplicates did not. Positive control was M. gallisepticum (1E+3 copies of target nucleic acid), and negative controls were water. Target capture and TMA was performed as is generally described herein. Amplification product was detected in real time using SEQ ID NO:14. In this example, none of the challenge organisms inhibited amplification of the target nucleic acid (M. gallisepticum). 1E+3 copies of M. gallisepticum was amplified in the presence of 1E+8 copies of each challenge organisms with average TTimes ranging from about 27.6 to about 28.8 minutes. Six of the challenge organisms were negative (0 of 4). S.epidermidis, E.coli; S. bovis and M. luteus challenge organisms each showed 1 of 4 positive in this experiment with TTimes of about 36.6 minutes, 42.4 minutes, 40.8 minutes and 56.5 minutes, respectively. L. pneumophilia challenge organism showed 3 of 4 positives in this experiment, with an average TTime of about 51.2 minutes (±13.7).

[190]. Example 6: Capture and Amplification of Mollecutes Nucleic Acid.

[191]. Target capture reagents were prepared to comprise the following: two target capture oligomers (SEQ ID NOS:28 & 29) and three tagged amplification oligomers (SEQ ID NOS:6, 30 & 31). The target capture oligomers are also blocker oligomers. The amplification reagent comprised a tag-targeting amplification oligomer (SEQ ID NO:33) and a promoter-based amplification oligomer (SEQ ID NO:15). An internal control was also used in these reactions (target capture reagent comprised SEQ ID NOS:32 & 35, amplification reagent comprised SEQ ID NOS:34 & 36). Amplification product was detected in real time using either SEQ ID NO:14 or SEQ ID NO:37. Target capture and amplification were performed as is generally described herein. Target nucleic acid was 1E+5 copies of *M. gallisepticum* 23S rRNA. Negative control was water. Each sample condition further comprised an *in vitro* transcript as internal control. Each test reaction condition was positive for 1E+5 copies of *M. gallisepticum*. There were no false positives.

- [192]. Example 7: Capture and Amplification of *Mollecutes* Nucleic Acid in the Presence of a Challenge Nucleic Acid.
- [193]. The performance of three different target capture systems was compared. Each system was tested with 1E+04 copies M. hominis (ATCC 41561) as positive control and C. perfringens (ATCC 13124) lysate as challenge nucleic acid at 1E+07 and 1E+08 copies. Three separate target capture reaction mixtures were prepared, each reaction mixture comprised four tagged amplification oligomers (SEQ ID NOS:5, 6, 7 & 8); and a first of the three reaction mixtures further comprised two target capture oligomers (SEO ID NOS:11 & SEQ ID NO:12) and two blocker oligomers (SEQ ID NOS:9 & 10), the second of the three reaction mixtures further comprised two target capture oligomers (SEQ ID NOS:64 & 65) and two blocker oligomers (SEQ ID NOS:9 & 10), and the third of the three reaction mixtures further comprised two target capture oligomers (SEQ ID NOS:28 & 29). In the third of the three reaction mixtures, the target capture oligomers were also used as blocker oligomers. The nucleotide sequences of the target hybridizing regions of both SEQ ID NOS:11 & 12 are 2-O-Me RNA residues. An internal control was also used, and thus all three reaction mixtures further comprised an in vitro transcript as internal control and SEQ ID NO:32 as a target capture oligomer for the in vitro transcript. Each of the target capture reagents was added to samples containing either the positive lysates or the cross reactor lysates. A negative control reaction comprised only water. These reactions were incubated at 45 minutes at 60.deg.C and then cooled at room temperature for 15 minutes. Target capture was then performed using magnetic beads and a KingFisher mL purification system.
- [194]. Following target capture, amplification reagent was added to each of the reaction conditions. The amplification reagent comprised a promoter-based amplification oligomer (SEQ ID NO:15) and two tag targeting oligomers, (SEQ ID NOS:13 & 33). Amplification oligomers directed to the internal control

were also included (SEQ ID NOS:34, 35 & 36). These reaction mixtures were then incubated at 42.deg.C, and amplification product was detected using one molecular torch for the sample and another for the internal control (SEQ ID NOS:14 & 37, respectively). Positive cut-off was set at 2000 RFU. Results are shown in Table 1. Sensitivity of each of the three target capture systems was comparable. The target capture oligomers comprising 2-O-Me nucleotide residues in the target capture regions were more selective against the challenge organism than were the deoxy target capture oligomers.

Target Capture Oligomer SEQ ID NOS:	Target / Target Amount	Total / # Positive	AVG TTime1 / SD	Avg TTime2 / SD
13 & 14	ATCC 13124/ 1E+7	12 / 3	35.8 / 8.92	39.36 / 1.79
13 & 14	ATCC 13124/ 1E+8	12 / 6	39.8 / 8.66	39.01 / 2.07
13 & 14	0 / 0	4 / 0	0 / 0	40.03 /
13 & 14	ATCC 41561 / 1E+4	4 / 4	24.0 / 0.86	44.32 /
11 & 12	ATCC 13124	12 / 0	0 / 0	36.30 / 2.82
11 & 12	ATCC 13124	12 / 1	40.7 / 0	35.99 / 2.61
11 & 12	0 / 0	4 / 0	0 / 0	35.17 / 0.78
11 & 12	ATCC 41561 / 1E+4	4 / 4	25.8 / 0.53	38.28 / 2.52
15 & 16	ATCC 13124	12 / 2	41.3 / 11.92	38.17 / 1.92
15 & 16	ATCC 13124	12 / 7	39.2 / 7.81	38.18 / 1.55
15 & 16	0 / 0	4 / 0	0 / 0	36.73 / 1.44
15 & 16	ATCC 41561 / 1E+4	4 / 4	25.6 / 1.53	38.91 / 1.51

[195]. Table 1. Summary of Results.

[196]. Example 8: Capture and Amplification of *Mollecutes* Nucleic Acid in the Presence of a Challenge Nucleic Acid.

[197]. The performance of two different target capture systems was compared using 1E+3 copies of *M. gallisepticum* as a positive control and using 1E+7 copies of *S. aureus* or ~2E+8 copies of *E. faecalis* as challenge organisms. In this example, the two lysis reaction mixtures were prepared as described in Example 7, above, for the first reaction mixture comprising two target capture oligomers (SEQ ID NOS:11 & SEQ ID NO:12) and two blocker oligomers (SEQ ID NOS:9 & 10), and for the third reaction mixture comprising two target capture oligomers (SEQ ID NOS:28 & 29). An internal control was used as described above. Each of the lysis reagents was added to samples containing either the positive lysates or

one of the cross reactor lysates. A negative control reaction comprised only water. These reactions were incubated at 45 minutes at 60.deg.C and then cooled at room temperature for 15 minutes. Target capture was then performed using magnetic beads and a KingFisher mL purification system. Amplification reagent, described above, was then added to each of the reaction conditions and the reactions were then incubated at 42.deg.C. A molecular torch was also present in the reaction for real time detection of amplification product(SEQ ID NOS:14 for the sample & 37 for the internal control). Positive cut-off was set at 2000 RFU.

- [198]. The target capture system comprising SEQ ID NOS:28 & 29 as target capture oligomers and SEQ ID NO:33 as a tag-targeting oligomer showed positive results for three of four samples containing 1E+3 copies of *M. gallisepticum*, with an average TTime of about 32.2 minutes (±12.85). This system provided no false positives and no cross reactivity with the *S. aureus* cross reacting organism. However, this system did show 6 of 12 positive reactions for *E. faecalis* with an average TTime of about 38.7 minutes (±7.18). Using the SEQ ID NO:13 tag-targeting oligomer with these same target capture oligomers (SEQ ID NOS:28 & 29) provided positive results for three of four samples containing 1E+3 copies of *M. gallisepticum*, with an average TTime of about 26 minutes (±2.76). This system provided no false positives and no cross reactivity with the *S. aureus* cross reacting organism. But, this system did show 7 of 12 positive reactions for *E. faecalis* with an average TTime of about 40.7 minutes (±4.9). The target capture system comprising SEQ ID NOS:11 & 12 as target capture oligomers and SEQ ID NO:13 as a tagtargeting oligomer showed provided positive results for two of four samples containing 1E+3 copies of *M. gallisepticum*, though one was only weakly positive, with an average TTime of about 41 minutes (±19.47). This system provided no false positives, no cross reactivity with the *S. aureus* challenge organism, and 1 of 12 positive reactions for *E. faecalis* with a TTime of about 37.8 minutes.
- [199]. Example 9: Amplification of Target Nucleic Acid from Various Species in the Class *Mollicutes* in the Presence of Challenge Nucleic Acids; Sensitivity in Multiplex Reactions.
- [200]. A multiplex reaction was performed to detect each of six different target nucleic acids in the presence of each of four challenge organisms. For this reaction, target organisms were *M. salvarium*, *M. hyorhinis*, *M. synoviae*, *M. arthritidis*, *M. pneumoniae* and *S. citri*. Challenge organisms were *B. subtilis*, *M. luteus*, *P. acnes* and *P. aeruginosa*. Positive control was *M. gallisepticum* and negative control was water. Target and challenge organisms were present at about 1E+4 copies (~1E+3 copies per CFU). Challenge organisms were also tested at about 1E+7 copies per reaction. Target capture reagent comprised the following: two target capture oligomers (SEQ ID NOS:11 & 12), four tagged amplification oligomers (SEQ ID NOS:5, 6, 7 & 8) and two blocker oligomers (SEQ ID NOS:9 & 10). Amplification reagent

comprised the following: a tag-targeting oligomer (SEQ ID NOS:13) and a promoter-based amplification oligomer (SEQ ID NO:15). A molecular torch was used for real time detection of amplifications product (SEQ ID NO:14). An internal control system was included as described herein and comprising an *in vitro* transcript and SEQ ID NOS:32, 34, 35, 36 & 37. Target capture, amplification and real-time detection of amplification product were performed as is generally described herein. Results showed that the *M. gallisepticum* control was positive with TTimes of about 20 to 23 minutes, though there was 1 of 8 false positives. All target organisms were amplified except for *M. pneumoniae*, which was later determined to have been the result of a miscalculation in concentration, see below. *M. salvarium* was positive with an average TTime of about 25 minutes, *M. hyorhinis* was positive with an average TTime of about 25 minutes, *M. synoviae* was positive with an average TTime of about 25 minutes, *M. pneumoniae* and *S. citri* was positive with an average TTime of about 25 minutes. Challenge organisms *B. subtilis, M. luteus, P. acnes* and *P. aeruginosa* were each mostly negative, though *B. subtilis* and *P. aeruginosa* had 1 of 8 positives and *M. luteus* showed a low riser (RFU greater than 2000 at a TTime of about 55-60 minutes).

[201]. An amplification reaction was performed as described above, but using a serial dilution of *M. pneumoniae* to determine whether the results obtained in the reaction directly above are due to a concentration error in the stock reagent or are attributable to the assay reagents. Target nucleic acids included 0 or 1E+4 copeis per reaction of the following: *M. hominis*, *A. laidlawii* and *S. citri*. A serial dilution was made for the stock *M. pneumoniae*, and a 1:10 and 1:100 dilution was tested. Positive control was *M. gallisepticum*. Negative control was water. Target capture, amplification and real-time detection of amplification product were performed as is generally described herein. Results showed an average TTime of about 24 minutes for *M. gallisepticum* positive control. *M. hominis* had an average TTime of about 23 minutes, *A. laidlawii* had an average TTime of about 27 minutes and *S. critri* had an average TTime of about 27 minutes. *M. pneomoniae* serial dilutions showed average TTimes of about 28 minutes for both the 1:10 dilution and the 1:100 dilution. There were no false positives.

[202]. Example 10: Exemplary Oligomers, Reference Sequences and Oligomer Regions.

[203]. In Table 2, below, there are exemplary oligomers, reference sequences and oligomer regions as described herein. The following nucleic acids are illustrated in this Table 2. Reference target nucleic acids are SEQ ID NOS:1-4. Tagged amplification oligomers, include SEQ ID NOS:5-8, 17-24, 30, 31, 40, 41, 49, 56, 57, 63 & 67-72. Tag closing regions (also called target closing regions) can be added or removed or modified on these tagged amplification oligomers as is described herein. Optionally, tag closing regions are joined to tag regions using one or more non-nucleotide linkers. Blocker oligomers include SEQ ID NOS:9, 10, 25 and 27. However, in some examples here, the target hybridizing region of a target capture

oligomer also served as a blocker oligomer. Target capture oligomers include SEQ ID NOS:11, 12, 28, 29, 64 & 65. Promoter-based oligomer includes SEQ ID NOS:15. Detection probes include SEQ ID NO:14 and SEQ ID NO:27. Tag-targeting amplification oligomers include SEQ ID NOS:13, 16, 33, 43, 44 & 66. These sequences or their complements, can also refer to tag region sequences contained within a tagged amplification oligomer, an initial amplification product and/or an amplification product. Target hybridizing regions include SEQ ID NOS:38, 39, 42, 46-48, 50-55 and 58-62. Internal control oligomers include SEQ ID NOS:32 & 34-37. SEQ ID NO:14 preferably comprises at least one 2'-O-Me residue, a 5' fluorescein and 3'dabcyl and/or a C9 linker between residues 20 and 21. SEQ ID NO:27 preferably comprises at least one 2'-O-Me residue, a 5' fluorescein and 3'dabcyl and/or a C9 linker between residues 5 and 6. Some oligomers further comprise one or more of a 2'-O-Me residue, a blocked 3'-terminus or a non-nucleotide linker. SEQ ID NO:77 is an exemplary promoter sequence for promoter-based oligomers.

[204]. Table 2: Exemplary Oligomers, Reference Sequences and Regions.

SEQ ID NO:	Sequence (5' to 3')
1	Mycoplasma hominis. GenBank Accession No. AF443616.3 GI:110815963. First seen at NCBI on July 28, 2006
2	Mycoplasma gallisepticum. GenBank Accession No. L08897.1 GI:530152. First seen at NCBI on August 17, 1994
3	Residues 75178 to 78004 of Acholeplasma laidlawii. CP000896.1 GI:161984995. The residues correspond to the 223S rRNA gene. GenBank Accession No. CP000896.1 GI:161984995 was first seen at NCBI on December 7, 2007.
4	Spiroplasma insolitum. GenBank Accession No. EU582532.1 GI:175362846. First seen at NCBI on April 12, 2008
5	CTAAATACTGACCTGCAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTATTTAG
6	TGTAACCATCTCCAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA
7	GCCGGcagtaggtaccgatgacgaGTAGCCGGCGTTTTCACCGGC
8	TGATCcagtaggtaccgatgacgaGCCGAGACAGCGAAGGGATCA
9	CGCCGUUCACUGGGGCUUC
10	CGCCGUUCACCCGGGCUUC
11	ACGUGUGUUCGUUCUCGGaaaaaaaaaaaaaaaaaaaaaa
12	CAACAGUUUUCUCGCGCGUtttaaaaaaaaaaaaaaaaaaaaaaa
13	TCGTcagtaggtaccgatgacga
14	ccAGGAAUUUCGCUACCUUAccugg
15	aatttaatacgactcactatagggagaACGGCCGTAACTATAACGGTCCTAAGG
16	gaacctagttgggcgagttacgga
17	gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTAGTTAG

18 CTAACTACTGGgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTAGTTAG 19 gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA 20 TACTGgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA 21 gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA 22 TGACCgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA 23 gaacctagttgggcgagttacggaCTTTCGTGCAGGTCA 24 GTAAATTCCGgaacctagttgggcgagttacggaCTTTCAAGCGGGACGGAATTTAC 25 CGCGUUCACCGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCGGGCUUCCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	SEQ ID NO:	Sequence (5' to 3')
TACTGGaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA 21 gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA 22 TGACCgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA 23 gaacctagttgggcgagttacggaTCACCATTCAAGCGGACGGAATTTAC 24 GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGACGGAATTTAC 25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGG 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaa 29 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaa	18	CTAACTACTGgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTAGTTAG
21 gaacctagttgggcgagttacggaCTTTCGTCCAGGTCA 22 TGACCgaacctagttgggcgagttacggaCTTTCGTCCAGGTCA 23 gaacctagttgggcgagttacggaTCATCGTCCAGGTCA 24 GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGAACGAATTTAC 25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaa	19	gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA
TGACCGaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA gaacctagttgggcgagttacggaTACACCATTCAAGCGGACGGATTTAC 24 GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGACGGATTTAC 25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaa	20	TACTGgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCAGTA
gaacctagttgggcagttacggaTACACCATTCAAGCGGACGGATTTAC 24 GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGACGGATTTAC 25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGGCUUC 27 gcgaaCGACAAGGAADUUCGC 28 CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaa 29 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaa 30 CTAATTACTGACCTGcagtaggtacgatgacgaTCGTGCAGGTCAGTAATTAG 31 AACAGACTCGGTcagtaggtacgatgacgaCAGCTTGATTTCACCGAGTCTGTT 32 CGUUCACUAUUGGUCUCUGCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaa 33 cagtaggtaccgatgacga 34 aatttaatacgactcactatagggagaccacaaAATAGATTATATAGGACGACAAGTAAAA 35 CUAUUGUCACUUCCUUGAGUAU 36 GACCATGTCCCAATTCGCACCAGG 37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgaTCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGGCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG	21	gaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA
24 GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGACGAATTTAC 25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	22	TGACCgaacctagttgggcgagttacggaCCTTTCGTGCAGGTCA
25 CGCCGUUCACCGGG 26 CGCCGUUCACCGGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaa 29 CGCCGUUCACCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaa 30 CTAATTACTGACCTGGagtaggtaccgatgaggaTCGTGCAGGTCAGTAATTAG 31 AACAGACTCGGTcagtaggtaccgatgaggaCAGCTTGATTTCACCGAGTCTGTT 32 CGUUCACUAUUGGUCUCCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaa	23	gaacctagttgggcgagttacggaTACACCATTCAAGCGGGACGGAATTTAC
26 CGCCGUUCACCGGGGCUUC 27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	24	GTAAATTCCGgaacctagttgggcgagttacggaTACACCATTCAAGCGGGACGGAATTTAC
27 gcgaaCGACAAGGAAUUUCGC 28 CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	25	CGCCGUUCACCGGG
CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	26	CGCCGUUCACCGGGGCUUC
CGCCGUUCACUGGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa	27	gcgaaCGACAAGGAAUUUCGC
CTAATTACTGACCTGcagtaggtaccgatgacgatCAGCTCAGTAATTAG AACAGACTCGGTcagtaggtaccgatgacgaCAGCTTGATTTCACCGAGTCTGTT CGUUCACUAUUGGUCUCUGCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa	28	CGCCGUUCACCCGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa
AACAGACTCGGTcagtaggtaccgatgacgaCAGCTTGATTTCACCGAGTCTGTT 32 CGUUCACUAUUGGCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa	29	CGCCGUUCACUGGGGCUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaa
CGUUCACUAUUGGUCUCUGCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa	30	CTAATTACTGACCTGcagtaggtaccgatgacgaTCGTGCAGGTCAGTAATTAG
aatttaatacgactcactatagggagaccacaaAATAGATTATATAGGACGACAAGTAAAA 35 CUAUUGUCACUUCCUUGAGUAU 36 GACCATGTCCCAATTCGCACCAGG 37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgaTCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGGCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGACGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGGT 49 CAGTAGGTACCGATGACGAGACAGTCAAGAGATGGTTACA	31	AACAGACTCGGTcagtaggtaccgatgacgaCAGCTTGATTTCACCGAGTCTGTT
aatttaatacgactcactatagggagaccacaaAATAGATTATATAGGACGACAAGTAAAA 35 CUAUUGUCACUUCCUUGAGUAU 36 GACCATGTCCCAATTCGCACCAGG 37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgatCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGCCGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGGTGAAGAGAGATGGTTACA		CGUUCACUAUUGGUCUCUGCAUUCtttaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa
35 CUAUUGUCACUUCCUUGAGUAU 36 GACCATGTCCCAATTCGCACCAGG 37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 39 CCTTTCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgatCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGCCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGACAGTCAAGAGATGGTTACA	33	cagtaggtaccgatgacga
36 GACCATGTCCCAATTCGCACCAGG 37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 39 CCTTTCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgatCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGCCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGGT 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	34	aatttaatacgactcactatagggagaccacaaAATAGATTATATAGGACGACAAGTAAAA
37 CCACUUGCGAUGUUUUAAGUGG 38 TCGTGCAGGTCAGTADTTAG 39 CCTTTCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgaTCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGGCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGGTTA 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA		CUAUUGUCACUUCCUUGAGUAU
TCGTGCAGGTCAGTADTTAG CCTTTCGTGCAGGTCAGTADTTAG CagtaggtaccgatgacgatCGTGCAGGTCAGTADTTAG CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG ACGCGGGCCGTAACTATAACGGTCCTAAGG Cagtaggtaccgatgacga Cagtaggtaccgatgacga Gagacctagttgggcgagttacgga TACACCATTCAAGCGGGACGGAATTTAC GAGACAGTCAAGAGATGGTTACA GAGACAGTCAAGAGATGGTTA AS GAGACAGTCAAGAGATGGTTA CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	36	GACCATGTCCCAATTCGCACCAGG
39 CCTTTCGTGCAGGTCAGTADTTAG 40 cagtaggtaccgatgacgatCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGCCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	37	CCACUUGCGAUGUUUUAAGUGG
40 cagtaggtaccgatgacgaTCGTGCAGGTCAGTADTTAG 41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGCCGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGACAGTCAAGAGATGGTTACA	38	TCGTGCAGGTCAGTADTTAG
41 CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG 42 ACGGCGGCCGTAACTATAACGGTCCTAAGG 43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	39	CCTTTCGTGCAGGTCAGTADTTAG
42 ACGGCGCCGTAACTATAACGGTCCTAAGG 43 Cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	40	cagtaggtaccgatgacgaTCGTGCAGGTCAGTADTTAG
43 cagtaggtaccgatgacga 44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	41	CAGTAGGTACCGATGACGACCTTTCGTGCAGGTCAGTADTTAG
44 gaacctagttgggcgagttacgga 45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	42	ACGGCGGCCGTAACTATAACGGTCCTAAGG
45 TACACCATTCAAGCGGGACGGAATTTAC 46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGAGATGGTTACA	43	cagtaggtaccgatgacga
46 GAGACAGTCAAGAGATGGTTACA 47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	44	gaacctagttgggcgagttacgga
47 GAGACAGTCAAGAGATGGTTA 48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	45	TACACCATTCAAGCGGGACGGAATTTAC
48 GAGACAGTCAAGAGATGG 49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	46	GAGACAGTCAAGAGATGGTTACA
49 CAGTAGGTACCGATGACGAGAGACAGTCAAGAGATGGTTACA	47	GAGACAGTCAAGAGATGGTTA
or a state of the	48	GAGACAGTCAAGAGATGG
50 CAGCTTGATTTCACCGAGTCTGTT	49	CAGTAGGTACCGATGACAGAGACAGTCAAGAGATGGTTACA
	50	CAGCTTGATTTCACCGAGTCTGTT

SEQ ID NO:	Sequence (5' to 3')
51	GGCGTTTTCACCGGCAGCT
52	AGCCGGCGTTTTCACCGGCAG
53	GGTAGCCGGCGTTTTCACCGGCAG
54	GTAGCCGGCGTTTTCACCGGC
55	TAATCGCGGGTAGCCGGCGTT
56	cagtaggtaccgatgacGCTTGATTTCACCGAGTCTGTT
57	cagtaggtaccgatgacgaGTAGCCGGCGTTTTCACCGGC
58	GCCGAGACAGCGAAGGGATCA
59	GCAGCCGAGACAGCGAAGGG
60	AATTTCACCGAGTCTGCAGCC
61	AAATTTCACCGAGTCTGCAGCC
62	CTAAAATTTCACCGAGTCTGCAGCC
63	cagtaggtaccgatgacgaGCCGAGACAGCGAAGGGATCA
64	CAACAGTTTTCTCGCGCGTTTTAAAAAAAAAAAAAAAAA
65	ACGTGTGTTCGCTTTAAAAAAAAAAAAAAAAAAAAAAAA
66	GTTGACGTACCGTATTGA
67	gttgacgtaccgtattgaGAGACAGTCAAGAGATGGTTACA
68	gttgacgtaccgtattgaTCGTGCAGGTCAGTAATTAG
69	gttgacgtaccgtattgaCAGCTTGATTTCACCGAGTCTGTT
70	AGCTGCCGGTGAcagtaggtaccgatgacgaGGCGTTTTCACCGGCAGCT
71	TCGACGCCACTcagtaggtaccgatgacgaGTAGCCGGCGTTTTCACCGGC
72	GTGAAATTTTAGcagtaggtaccgatgacgaCTAAAATTTCACCGAGTCTGCAGCC
73	GTTGACGTACCGTATTGACCTTTCGTGCAGGTCAGTAGTTAG
74	GTTGACGTACCGTATTGATTTCGTGCAGGTCAGTAGTTAG
75	GTTGACGTACCGTATTGATCGTGCAGGTCAGTAGTTAG
76	GTTGACGTACCGTATTGAGTGCAGGTCAGTAGTTAG
78	CGCCGUUCACYKGGGCUUC

[205]. The methods illustratively described herein may suitably be practiced in the absence of any element or elements, limitation or limitations, not specifically disclosed herein. Thus, for example, the terms "comprising", "including," containing", etc. shall be read expansively and without limitation. Additionally, the terms and expressions employed herein have been used as terms of description and not of limitation, and there is no intention in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof. It is recognized that various modifications are

possible within the scope of the invention claimed. Thus, it should be understood that although the present invention has been specifically disclosed by preferred embodiments and optional features, modification and variation of the invention embodied therein herein disclosed may be resorted to by those skilled in the art, and that such modifications and variations are considered to be within the scope of this invention.

[206]. The invention has been described broadly and generically herein. Each of the narrower species and subgeneric groupings falling within the generic disclosure also form part of the methods. This includes the generic description of the methods with a proviso or negative limitation removing any subject matter from the genus, regardless of whether or not the excised material is specifically recited herein.

[207]. Other embodiments also fall within the following claims. In addition, where features or aspects of the methods are described in terms of Markush groups, those skilled in the art will recognize that the invention is also thereby described in terms of any individual member or subgroup of members of the Markush group.

CLAIMS:

A method for the in vitro amplification of a nucleic acid in a sample, said nucleic acid being from one or more species in the class Mollicutes, comprising the steps of: (a.) contacting a sample with at least two tagged amplification oligomers, each of which individually comprises a target hybridizing region and a tag region, wherein said at least two tagged amplification oligomers are selected from the group consisting of: (i) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; (ii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; (iii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and (iv) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4; and (b.) providing suitable conditions for performing an in vitro amplification reaction.

- 2. The method of claim 1, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 24 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1.
- 3. The method of claim 2, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:38 and SEQ ID NO:39.
- 4. The method of claim 2, wherein said tag region is from 19 to 24 nucleobases in length.
- 5. The method of claim 4, wherein said tag region has a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; SEQ ID NO:43, the complement of SEQ ID NO:43, SEQ ID NO:44 and the complement of SEQ ID NO:44.
- 6. The method of claim 5, wherein said tagged amplification oligomer is a sequence selected from the

group consisting of: SEQ ID NO:17, SEQ ID NO:40 and SEQ ID NO:41.

7. The method of claim 2, claim 4 or claim 5, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into the closed inactive configuration.

- 8. The method of claim 7, wherein said tagged amplification oligomer has at least 95% sequence identity to a nucleotide sequence selected from the group consisting of: SEQ ID NO:5, SEQ ID NO:18 and SEQ ID NO:30.
- 9. The method of claim 7, wherein said tagged amplification oligomer is SEQ ID NO:5.
- 10. The method of claim 1, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 18 to 28 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2.
- 11. The method of claim 10, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:45, SEQ ID NO:46, SEQ ID NO:47 and SEQ ID NO:48.
- 12. The method of claim 10, wherein said tag region is from 18 to 24 nucleobases in length.
- 13. The method of claim 12, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; SEQ ID NO:43, the complement of SEQ ID NO:43; SEQ ID NO:44, and the complement of SEQ ID NO:44.
- 14. The method of claim 13, wherein said tagged amplification oligomer is a nucleotide sequence selected from the group consisting of: SEQ ID NO:23 and SEQ ID NO:49.
- 15. The method of claim 10, claim 12 or claim 13, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into a closed inactive configuration.

16. The method of claim 15, wherein said tagged amplification oligomer has at least 95% sequence identity to a nucleotide sequence selected from the group consisting of: SEQ ID NO:6 and SEQ ID NO:24.

- 17. The method of claim 15, wherein said tagged amplification oligomer is SEQ ID NO:6.
- 18. The method of claim 1, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 19 to 24 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3.
- 19. The method of claim 18, wherein said target hybridizing region is sequence selected from the group consisting of: SEQ ID NO:50, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:54 and SEQ ID NO:55.
- 20. The method of claim 18, wherein said tag region is from 18 to 24 nucleobases in length.
- 21. The method of claim 20, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; SEQ ID NO:43, SEQ ID NO:44, the complement of SEQ ID NO:43, and the complement of SEQ ID NO:44.
- 22. The method of claim 18, claim 19 or claim 21, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into the closed inactive configuration.
- 23. The method of claim 22, wherein said tagged amplification oligomer has at least 95% sequence identity to a nucleotide sequence selected from the group consisting of: SEQ ID NO:7 and SEQ ID NO:31.
- 24. The method of claim 22, wherein said tagged amplification oligomer is SEQ ID NO:7.
- 25. The method of claim 1, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 25 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.

26. The method of claim 25, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 and SEQ ID NO:62.

- 27 The method of claim 25, wherein said tag region is from 18 to 24 nucleobases in length.
- 28. The method of claim 27, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43, SEQ ID NO:44, the complement of SEQ ID NO:43, and the complement of SEQ ID NO:44.
- 29. The method of claim 25, claim 26 or claim 28, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into the closed inactive configuration.
- 30. The method of claim 29, wherein said tagged amplification oligomer has at least 95% sequence identity to SEQ ID NO:8.
- 31. The method of claim 29, wherein said tagged amplification oligomer is SEQ ID NO:8.
- 32. The method of claim 1, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:5 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer consisting of SEQ ID NO:6, a tagged amplification oligomer consisting of SEQ ID NO:8.
- 33. The method of claim 1, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:6 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer consisting of SEQ ID NO:5, a

tagged amplification oligomer consisting of SEQ ID NO:7 and a tagged amplification oligomer consisting of SEQ ID NO:8.

- 34. The method of claim 1, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:7 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer consisting of SEQ ID NO:5, a tagged amplification oligomer consisting of SEQ ID NO:6 and a tagged amplification oligomer consisting of SEQ ID NO:8.
- 35. The method of claim 1, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:8 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer consisting of SEQ ID NO:5, a tagged amplification oligomer consisting of SEQ ID NO:5, a tagged amplification oligomer consisting of SEQ ID NO:7.
- 36. The method of any one of claims 32, 33, 34 or 35, further comprising a blocker oligomer consisting of SEQ ID NO:78.
- The method of claim 1, wherein said at least two tagged amplification oligomers comprises four tagged amplification oligomers selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer consisting of SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer consisting of SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer consisting of SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, and a tagged amplification oligomer consisting of SEQ ID NO:8.
- 38. The method of claim 37, further comprising at least one blocker oligomer selected from the group consisting of: SEQ ID NO:9, SEQ ID NO:78 and SEQ ID NO:10.
- 39. The method of claim 1, further comprising at least one blocker oligomer.

40. The method of claim 39, wherein said at least one blocker oligomer is selected from the group consisting of: SEQ ID NO:9, SEQ ID NO:78 and SEQ ID NO:10.

- 41. The method of any one of claims 1 through 49, wherein the amplification reaction of step (b.) generates a first amplification product containing a tag sequence, said tag sequence being selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- 42. The method of any one of claims 1 through 41 wherein the amplification reaction of step (b.) is an isothermal amplification reaction.
- 43. The method of any one of claims 1 through 42, further comprising the steps of (c.) providing at least one additional amplification oligomer selected from the group consisting of a tag-targeting amplification oligomer, a primer oligomer and a promoter-based amplification oligomer; and (d.) providing suitable conditions for performing a second amplification reaction using the at least one amplification oligomer from step (c.) and using the initial amplification product generated in step (b.) as a target nucleic acid.
- 44. The method of claim 43, wherein said additional amplification oligomer is a tag-targeting amplification oligomer comprising a target hybridizing region that contains a nucleotide sequence configured to specifically hybridize to a tag sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- 45. The method of claim 44, wherein said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length.
- 46. The method of claim 44, wherein said tag-targeting amplification oligomer has a sequence that is at least 95% identical to a sequence selected from the group consisting of: SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.
- 47. The method of claim 46, wherein said tag-targeting amplification oligomer is SEQ ID NO:13.
- 48. The method of claim 43, wherein said additional amplification oligomer is a promoter-based

amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1.

- 49. The method of claim 48, wherein said target hybridizing region of said promoter-based amplification oligomer has a sequence that is at least 95% identical to SEQ ID NO:42.
- 50. The method of claim 48, wherein said target hybridizing region of said promoter-based amplification oligomer is a sequence consisting of SEO ID NO:42.
- 51. The method of claim 48, wherein said promoter-based amplification oligomer is SEQ ID NO:15.
- 52. The method of claim 41, wherein said at least one additional amplification oligomer comprises a first amplification oligomer that is a tag-targeting amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:13, and a second amplification oligomer that is a promoter-based amplification oligomer with a nucleotide sequence that is at least 95% identical to SEQ ID NO:15.
- 53. The method of claim 52, wherein said tag targeting oligomer is SEQ ID NO:13.
- 54. The method of claim 53, wherein said promoter-based amplification oligomer is SEQ ID NO:15.
- 55. The method of any one of claims 43 through 54, wherein the amplification reaction of step (d.) is an isothermal amplification reaction.
- 56. The method of any one of claims 1 through 55, further comprising a detection step to identify an initial amplification product generated in step (b.), an amplification product generated in step (d.) or the amplification products generated in steps (b.) and (d.).
- 57. The method of claim 56, wherein said detection step is a probe-based detection step.
- 58. The method of claim 57, comprising a detection probe that is SEQ ID NO:14.
- 59. The method of any of claims 1 through 58, further comprising a target capture step using at least one target capture oligomer.
- 60. The method of claim 59, wherein at least one of said target capture oligomer is also a blocker oligomer.
- 61. The method of claim 59, wherein said at least one target capture oligomer is selected from the

group consisting of SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:28, SEQ ID NO:29, SEQ ID NO:64, SEQ ID NO:65 and combinations thereof.

- 62. The method of any one of claims 1 through 61, wherein said species of the class *Mollicutes* is selected from the group consisting of: *Mycoplasma* species, *Acholeplasma* species, *Spiroplasma* species and combinations thereof.
- 63. The method of claim 7, 15, 22 or 29, wherein said tag-closing region is joined to said tag region by a non-nucleotide linker.
- 64. A reaction mixture for generating from one or more species of the class Mollicutes in a sample an initial nucleic acid amplification product containing a tag sequence or complement thereof, wherein said reaction mixture comprises at least two tagged amplification oligomers selected from the group consisting of: (i) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; (ii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; (iii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and (iv) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.
- 65. The reaction mixture of claim 64, wherein said reaction mixture further comprises a blocker oligomer consisting of SEQ ID NO:78.
- 66. The reaction mixture of claim 64, wherein said reaction mixture further comprises two blocker oligomers, the first consisting of SEQ ID NO:9 and the second consisting of SEQ ID NO:10.
- 67. The reaction mixture of claim 64, further comprising at least one target capture oligomer.
- 68. The reaction mixture of claim 67, wherein said at least one target capture oligomer is selected from the group consisting of SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:28, SEQ ID NO:29, SEQ ID NO:64, SEQ ID NO:65 and combinations thereof.

69. An amplification reaction mixture for generating a nucleic acid amplification product from an initial nucleic acid amplification product containing a tag sequence or complement thereof and that was generated from a species of the class *Mollicutes*, wherein said amplification reaction mixture comprises:

(a) a tag-targeting amplification oligomer comprising a target hybridizing region containing a nucleic acid sequence configured to hybridize to said tag sequence or complement thereof on said initial nucleic acid amplification product, and (b) a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1.

- 70. The amplification reaction mixture of claim 69, wherein said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42.
- 71. The amplification reaction mixture of claim 69, wherein said promoter-based amplification oligomer is SEQ ID NO:15.
- 72. The amplification reaction mixture of claim 69, wherein said target hybridizing region of said tagtargeting amplification oligomer is 18 to 24 nucleobases in length.
- 73. The amplification reaction mixture of claim 72, wherein said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:13.
- 74. The amplification reaction mixture of claim 69, wherein said tag-targeting amplification oligomer is SEQ ID NO:13.
- A composition for determining the presence or absence of one or more species from the Mollicutes class in a sample, said composition comprising at least two tagged amplification oligomers, each of which individually comprises a target hybridizing region and a tag region, wherein said at least two tagged amplification oligomers are selected from the group consisting of: (i) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1; (ii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 4752 to 4798 of SEQ ID NO:2; (iii) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a

nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3; and (iv) a tagged amplification oligomer comprising a target hybridizing region that is from 15 to 50 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.

- 76. The composition of claim 75, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 24 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5065 to 5088 of SEQ ID NO:1.
- 77. The composition of claim 76, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:38 and SEQ ID NO:39.
- 78. The composition of claim 76, wherein said tag region is 19 nucleobases in length.
- 79. The composition of claim 76, wherein said tag region is 24 nucleobases in length.
- 80. The composition of claim 76, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- 81. The composition of any one of claims 76 through 80, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into a closed inactive configuration.
- 82. The composition of claim 81, wherein said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:5.
- 83. The composition of claim 81, wherein said tagged amplification oligomer is SEQ ID NO:5.
- 84. The composition of claim 75, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 18 to 28 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid

corresponding to residues 4752 to 4798 of SEQ ID NO:2.

85. The composition of claim 84, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:45, SEQ ID NO:46, SEQ ID NO:47 and SEQ ID NO:48.

- 86. The composition of claim 84, wherein said tag region is from 18 to 24 nucleobases in length.
- 87. The composition of claim 84, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- 88. The composition of any one of claims 84 through 87, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into a closed inactive configuration.
- 89. The composition of claim 88, wherein said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:6.
- 90. The composition of claim 88, wherein said tagged amplification oligomer is SEO ID NO:6.
- 91. The composition of claim 75, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 19 to 24 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1954 to 2006 of SEQ ID NO:3.
- 92. The composition of claim 91, wherein said target hybridizing region is selected from the group consisting of: SEQ ID NO:50, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:54 and SEQ ID NO:55.
- 93. The composition of claim 92, wherein said tag region is from 10 to 25 nucleobases in length.
- 94. The composition of claim 92, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100%

identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.

- 95. The composition of any one of claims 91 through 94, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into the closed inactive configuration.
- 96. The composition of claim 95, wherein said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:7.
- 97. The composition of claim 95, wherein said tagged amplification oligomer is SEO ID NO:7.
- 98. The composition of claim 75, wherein one of said at least two tagged amplification oligomers comprises a target hybridizing region that is from 20 to 25 nucleobases in length and contains a nucleotide sequence that is configured to specifically hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 1994 to 2036 of SEQ ID NO:4.
- 99. The composition of claim 98, wherein said target hybridizing region is a sequence selected from the group consisting of: SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61 and SEQ ID NO:62.
- 100. The composition of claim 98, wherein said tag region contains a nucleotide sequence selected from the group consisting of: a sequence with at least 95% identity to SEQ ID NO:43 or complement thereof; a sequence with at least 95% identity to SEQ ID NO:44 or complement thereof; a sequence with 100% identity to SEQ ID NO:43 or complement thereof; and a sequence with 100% identity to SEQ ID NO:44 or complement thereof.
- 101. The composition of any one of claims 98 through 100, wherein said tagged amplification oligomer further comprises a tag-closing region that is 3 to 20 nucleobases in length and is substantially complementary to a portion of said target hybridizing region so as to hybridize under a set of conditions to all or a portion of said target hybridizing region and thereby place the tagged amplification oligomer into the closed inactive configuration.
- 102. The composition of claim 101, wherein said tagged amplification oligomer comprises at least 95% sequence identity to SEQ ID NO:8.

103. The composition of claim 101, wherein said tagged amplification oligomer is SEQ ID NO:8.

- 104. The composition of claim 75, further comprising at least one blocker oligomer.
- 105. The composition of claim 104, wherein one of said at least one blocker oligomer selected from the group consisting of: SEQ ID NO:78, SEQ ID NO:9 and SEQ ID NO:10.
- 106. The composition of claim 75, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:5 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:7 and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8.
- 107. The composition of claim 75, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:6 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:7 and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8.
- 108. The composition of claim 75, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:7 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8.
- 109. The composition of claim 75, wherein said at least two tagged amplification oligomers comprises a first tagged amplification oligomer with a nucleotide sequence consisting of SEQ ID NO:8 and a second tagged amplification oligomer selected from the group consisting of: a tagged amplification oligomer with

a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6 and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:7.

- 110. The composition of any one of claims 106 through 109, further comprising at least one blocker oligomer consisting of SEQ ID NO:78.
- 111. The composition of claim 75, wherein said at least two tagged amplification oligomers comprises four tagged amplification oligomers selected from the group consisting of: a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:5, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:6, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:7, a tagged amplification oligomer with a sequence that is at least 95% identical to SEQ ID NO:8, and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8, and a tagged amplification oligomer with a sequence that is 100% identical to SEQ ID NO:8.
- 112. The composition of claim 111, further comprising at least one blocker oligomer with a sequence selected from the group consisting of: SEQ ID NO:78, SEQ ID NO:9 and SEQ ID NO:10.
- 113. The composition of claim 81, 88, 95 or 101, wherein said tag-closing region is joined to said tag region by a non-nucleotide linker.
- 114. A kit comprising at least two tagged amplification oligomers and a tag targeting amplification oligomer, wherein said at least two tagged amplification oligomer are selected from the group of tagged amplification oligomers in any one of claims 75-113, and wherein said tag-targeting amplification oligomer comprises a target hybridizing region that contains a nucleotide sequence configured to hybridize to the tag region of each of said at least one tagged amplification oligomers or complements thereof.
- 115. The kit of claim 114, wherein said target hybridizing region of said tag-targeting amplification oligomer is 18 to 24 nucleobases in length.
- 116. The kit of claim 115, wherein said tag-targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NO:13, SEQ ID NO:16 and SEQ ID NO:33.

117. The kit of claim 115, wherein said tag-targeting amplification oligomer is SEQ ID NO:13.

- 118. The kit of claim 114, wherein each of said tagged amplification oligomers comprise a tag region that is at least 95% identical to SEQ ID NO:43.
- 119. The kit of claim 114, wherein each of said tagged amplification oligomers comprise a tag region that is at least 95% identical to SEQ ID NO:44
- 120. The kit of claim 114, wherein each of said tagged amplification oligomers comprise a tag region that is SEQ ID NO:44.
- 121. The kit of claim 114, wherein each of said tagged amplification oligomers comprise a tag region that is SEQ ID NO:43.
- 122. The kit of any one of claims 118 through 121, wherein said tag targeting amplification oligomer comprises a sequence that is at least 95% identical to a sequence selected from the group consisting of SEQ ID NOS:13, 16 and 33.
- 123. The kit of claim 121, wherein said tag targeting amplification oligomer is SEQ ID NO:13.
- 124. The kit of claim 114, further comprising a promoter-based amplification oligomer comprising a target hybridizing region that is 10 to 50 nucleobases in length and contains a nucleotide sequence that is configured to hybridize to all or a portion of a region of a target nucleic acid corresponding to residues 5016 to 5045 of SEQ ID NO:1.
- 125. The kit of claim 122, wherein said target hybridizing region of said promoter-based amplification oligomer comprises a sequence that is at least 95% identical to SEQ ID NO:42.
- 126. The kit of claim 122, further comprising a promoter-based amplification oligomer comprising a sequence that is at least 95% identical to SEQ ID NO:15.
- 127. The kit of claim 126, wherein said promoter-based amplification oligomer comprising a sequence that is 100% identical to SEQ ID NO:15.
- 128. The kit of any of claims 114 through 127, further comprising one or more of a target capture oligomer, a detection probe, a blocker or combinations thereof.

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Relevant to claim No.

1 - 128

a. classification of subject matter INV. C12Q1/68 ADD.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

According to International Patent Classification (IPC) or to both national classification and IPC

Category*

Υ

Minimum documentation searched (classification system followed by classification symbols) C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

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X Further documents are listed in the continuation of Box C.	X See patent family annex.	
 Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance 	"T" later document published after the inter or priority date and not in conflict with cited to understand the principle or the invention	the application but
"E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or	"X" document of particular relevance; the cl cannot be considered novel or cannot involve an inventive step when the doc	be considered to
which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means	"Y" document of particular relevance; the ci cannot be considered to involve an inv document is combined with one or mo ments, such combination being obviou	rentive step when the re other such docu–
"P" document published prior to the international filing date but later than the priority date claimed	in the art. "&" document member of the same patent f	amily
Date of the actual completion of the international search	Date of mailing of the international sear	ch report
18 August 2010	30/08/2010	
Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk	Authorized officer	
Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Reuter, Uwe	

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A	US 6 074 826 A (HOGAN JAMES J [US] ET AL) 13 June 2000 (2000-06-13) example 9; sequence 58	1–128
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1	DATABASE EMBL [Online] 26 September 2002 (2002-09-26), "Mycoplasma hominis DNA-dependent RNA polymerase alpha chain gene, partial cds; 50S ribosomal protein L17 and 50S ribosomal protein L32 genes, complete cds; rrnA operon, complete sequence; 50S ribosomal protein L10 and 50S ribosomal protein L7/L12 genes, complete cds; and DNA-dependent RNA polymerase" XP002597076 retrieved from EBI accession no. EMBL:AF443616 Database accession no. AF443616 the whole document	
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Вох	No. I	Nucleotide and/or amino acid sequence(s) (Continuation of item 1.b of the first sheet)
1.	With	regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed nation, the international search was carried out on the basis of:
	a.	(means) X on paper X in electronic form
2.	b.	(time) X in the international application as filed together with the international application in electronic form subsequently to this Authority for the purpose of search In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed
	Ш	or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
3.	Addit	ional comments:

Information on patent family members

International application No
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