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#### Abstract

**Background:** Ribosomal RNA molecules are widely used for phylogenetic and *in situ* identification of bacteria. Nevertheless, their use to distinguish microorganisms within a species is often restricted by the high degree of sequence conservation and limited probe accessibility to the target in fluorescence *in situ* hybridization (FISH). To overcome these limitations, we examined the use of tmRNA for *in situ* identification. In *E. coli*, this stable 363 nucleotides long RNA is encoded by the *ssrA* gene, which is involved in the degradation of truncated proteins.

**Results:** Conserved sequences at the 5'- and 3'-ends of tmRNA genes were used to design universal primers that could amplify the internal part of ssrA from Gram-positive bacteria having low G+C content, i.e. genera *Bacillus, Enterococcus, Lactococcus, Lactobacillus, Leuconostoc, Listeria, Streptococcus* and *Staphylococcus*. Sequence analysis of tmRNAs showed that this molecule can be used for phylogenetic assignment of bacteria. Compared to 16S rRNA, the tmRNA nucleotide sequences of some bacteria, for example *Listeria*, display considerable divergence between species. Using *E. coli* as an example, we have shown that bacteria can be specifically visualized by FISH with tmRNA targeted probes.

**Conclusions:** Features of tmRNA, including its presence in phylogenetically distant bacteria, conserved regions at gene extremities and a potential to serve as target for FISH, make this molecule a possible candidate for identification of bacteria.

#### Background

In recent years, molecular approaches based on nucleotide sequences of ribosomal RNA (rRNA) have become widely used tools for identification of bacteria [1–4]. The high degree of evolutionary conservation makes 16S and 23S rRNA molecules very suitable for phylogenetic studies above the species level [3–5]. More than 16,000 sequences of 16S rRNA are presently available in public databases [4,6]. The 16S rRNA sequences are commonly used to design fluorescently labeled oligonucleotide probes. Fluorescence *in situ* hybridization (FISH) with these probes followed by observation with epifluores-

Received: 21 May 2001 Accepted: 7 September 2001 cence microscopy allows the identification of a specific microorganism in a mixture with other bacteria [2–4]. By shifting probe target sites from conservative to increasingly variable regions of rRNA, it is possible to adjust the probe specificity from kingdom to species level.

Nevertheless, 16S rRNA sequences of closely related strains, subspecies, or even of different species are often identical and therefore can not be used as differentiating markers [3]. Another restriction concerns the accessibility of target sites to the probe in FISH experiments. The presence of secondary structures, or protection of rRNA segments by ribosomal proteins in fixed cells can limit the choice of variable regions as *in situ* targets for oligonucleotide probes [7,8]. One way to overcome the limitations of in situ identification of bacteria is to use molecules other than rRNA for phylogenetic identification of bacteria, for which nucleotide sequences would be sufficiently divergent to design species specific probes, and which would be more accessible to oligonucleotide probes. For this purpose we investigated the possibility of using tmRNA (also known as 10Sa RNA; [9–11]). This molecule was discovered in *E. coli* and described as small stable RNA, present at ~1,000 copies per cell [9,11]. The high copy number is an important prerequisite for FISH, which works best with naturally amplified target molecules. In E. coli, tmRNA is encoded by the ssrA gene, is 363 nucleotides long and has properties of tRNA and mRNA [12,13]. tmRNA was shown to be involved in the degradation of truncated proteins: the tmRNA associates with ribosomes stalled on mRNAs lacking stop codons, finally resulting in the addition of a C-terminal peptide tag to the truncated protein. The peptide tag directs the abnormal protein to proteolysis [14,15]. 165 tmRNA sequences have so far (August 2001; The tmRNA Website: [http://www.indiana.edu/~tmrna/]) been determined [16,17]. The tmRNA is likely to be present in all bacteria and has also been found in algae chloroplasts, the cyanelle of Cyanophora paradoxa and the mitochondrion of the flagellate Reclinomonas americana [10,17,18].

In this study we present the first data on suitability of tm-RNA for phylogenetic affiliation and *in situ* identification of bacteria.

## Results and discussion Analysis of ssrA sequences from the species Lactococcus lactis

*Lactococcus lactis* was chosen to investigate the variability of tmRNA within one species. We used oligonucleotides B1 and B2, designed according to conserved tmRNA regions of *B. subtilis* (Table 1; [13]) to PCR-amplify the internal part of the *ssrA* gene from *L. lactis* strain IL1403 (named by analogy with the *E. coli* tmRNA gene; [12]). The nucleotide sequence of the amplified fragment was determined using the same primers. In-

tactgacaga	aaagttaaaa	tatgctataa	ttagacttca	
aggttcaaag	aattactgac <u>11</u>	agaataatca L	gtaggtatct —>	
ccttaaattt	GGGGGCGTTA	CGGATTCGAC	AGGCATTGTC	30
GCGCATGAAC	TGCAACTGCT	GAGGGATCAG	GATAATCATC	70
CGCAGATAAA	A TATAACT <i>GCT</i>	к п п ААААА <i>Т</i> ААТА	T Q T Y CACAAACTTA	110
A M A CGCAATGGCA	a stop GCCTAAACAG	CACCATGCGT	GCCTGATTTT	150
TGCTC <b>T</b> CTGA	TGGCAATTTG 208	ACGGCCTAAA 4L	CTTTTAGTCA 3L	190
GATACGTTGT	TGTGGAG <b>G</b> CT	TGACGCAGCA	AAAGAGATTT	230
 AAAGCCCCGC	AAAATGTCGC	TGTTTGAGAC	TGGCTTTGGC	270
ATTTTGTTAA	ATTTGAGAAA	GTCCTATGGT	TGTAGACGTT 12L	310
GATGTAGCAA	GGTGTTTGGA	CAGGGGTTCG	ACTCCCCTCG	350
CCTCCAttga	atggatttta	tagaatccaa	tagtattaga	
aatcgctcaa	tagagcggtt	tttcttttct	ctagtccgat	
atgatacgat	acaatctgat	aaaagggagg	gagaaaaaga	
gggagaaaaa	ataaaaaaac	cagtttatta		

#### Figure I

Nucleotide sequence of the *L. lactis* IL1403 chromosomal fragment encoding the *ssrA* gene. In capital letters, *ssrA* gene; in italics, presumed tag sequence, followed by a stop codon, the putative peptide is shown above the DNA sequence; in bold, variable positions 156, 208 and 235; underlined, regulatory regions (-10, -35 and terminator). Localization of primers 3L, 4L, 11L and 12L indicated by arrows.

verse PCR was used to obtain the sequence of the complete gene and of flanking regions [19]. For that purpose, chromosomal DNA of IL1403 was digested separately with restriction endonucleases *BclI*, *BglII*, *BspHI*, *HindIII*, *NsiI* or *SpeI*, respectively. The digested lactococcal DNA from each restriction reaction was ligated and PCRamplified, using primers L3 and L4 (Fig. 1, Table 1). A PCR fragment of approximately 2,000 bp was obtained from *Hind*III digested and religated DNA. The fragment was sequenced using primers L3, L4, L11 and L12 (Table 1) to obtain the full sequence of *ssrA* and surrounding regions (Fig. 1). The limits of the lactococcal *ssrA* gene were determined after alignment of this sequence with that of the *B. subtilis ssrA* gene [13].

The lactococcal tmRNA gene is 356 bp long and surrounded by typical regulatory sequences, suggesting a monocistronic operon structure (Fig. 1). The terminal 3'-CCA triplet of tmRNA is not encoded by the gene, and may be added posttrancriptionally, as in the case of *B. subtilis*[13]. The conservative peptide tag sequence, which is between UAA triplets may be distinguished in the central part of the gene. This sequence allows to pre-

Oligonucleotide	Sequence 5'-3' *	Target tmRNA
BI	ggggacgttacggattcgac, 1-20	B. subtilis
B2	tggagacgccgggagtcgaa, 351–360	B. subtilis
L3	ttaaagccccgcaaaatgtc, 229–248	L. lactis IL1403
L4	gtcaagcctccacaacaacg, 195–214	L. lactis IL1403
LII	acggattcgacagg, 10–23	L. lactis IL1403
LI2	gggagtcgaaccc, 234–246	L. lactis IL1403
EI	F-gtggaatccagaatcagcccc, 1–21	E. coli
E2	F-gcgtagttttcgtc, 96–109	E. coli
E3	F-ttgatccccgtcctaagagcgg, 154–175	E. coli
E4	F-tctcttttgggtttgacctctc, 176–197	E. coli
E5	F-tggtggagctggcgggagttgaa, 341–363	E. coli
RNAI	HRP-tgggtttgacctctcttg, 173–190	E. coli

#### Table 1: Oligonucleotides used in this work.

\* The type of label is either fluorescein (F-) or horseradish peroxidase (HRP-). Target sites are given after the sequences as nucleotide position numbers of the corresponding tmRNAs.

dict a 12 amino acid peptide (AKNNTQTYAMAA; see Fig. 1), which resembles the *ssrA*-encoded *E. coli* tmRNA tag sequence (ANDENYALAA; [14,15]). These similar features indicate that in Gram-positive bacteria, as in *E. coli*, the *ssrA* gene product may be involved in the turnover of truncated proteins.

To design universal primers for PCR amplification of tm-RNA genes from different Gram-positive bacteria, we aligned tmRNA sequences of L. lactis and B. subtilis. The primers 11L and 12L corresponded to the conserved 5'and 3' ends of the tmRNA molecule, which are identical in L. lactis and B. subtilis (Fig. 1). These primers were used for PCR amplification and sequencing of internal segments of ssrA from different L. lactis strains. Alignment of tmRNA sequences of 16 L. lactis strains showed that there were three variable positions in the gene (156, 208 and 235, Fig. 1). These differences between L. lactis tmRNA sequences allowed us to distinguish 3 groups within the species, which corresponded to L. lactis ssp. lactis, L. lactis ssp. lactis biovar.diacetylactis and L. lactis ssp. cremoris (Table 1). Strain IL890 is an exception, since by ssrA sequence it belongs to the L. lactis ssp. lactis group, although it is classified as biovar. diacetylactis. IL2961 differs from the other *L. lactis* ssp. *lactis* strains by an T in the variable position 235 and a C in position 167 (Fig. 1). These differences of IL2961 are not surprising since this strain has an unusual RAPD pattern [20]. Even if the number of strains tested and the divergence of lactococcal tmRNA sequences are too low to make a significant clustering, these results show that biovariety diacetylactis may be distinguished by differences in chromosomal sequences not related to diacetyl production or to a citrate permease encoding plasmid, which is in line with previous reports [21–23]. Comparison of tm-RNA sequences from a larger collection of lactococcal strains will allow us to confirm this finding.

# Phylogenetic analysis of ssrA genes from gram-positive bacteria

We PCR-amplified and sequenced the internal part of the ssrA gene of 24 strains of Gram-positive bacteria with low G+C content, namely of the genera Bacillus, Lactococcus, Lactobacillus, Streptococcus, Staphylococcus, Leuconostoc, Listeria and Enterococcus. The divergence of tmRNA sequences within one genus differed according the species. For example, we found 5 base differences between ssrA genes of Listeria innocua and Listeria monocytogenes (1.3% of divergence). The 16S rRNA sequences of these species are less divergent (0.5%). In staphylococci, S. saprophyticus tmRNA differs from S. xylosus in only one base (0.3% of divergence). In contrast, the difference between these two species and S. epidermidis is more marked (more than 20% divergence). The divergence between species within the genera Lactococcus, Lactobacillus, Leuconostoc and Enterococcus is 8.9%, 2.9%, 18% and 6.6%, respectively. Sequence comparisons showed that in some cases, as for Listeria innocua and Listeria monocytogenes, the tmR-NAs are more divergent than 16S RNA, thus making tm-RNA a promising tool for species identification. Although the total number of base differences even in a less divergent 16S rRNA sequence might be higher or at least equal to that of the corresponding tmRNA sequence, the advantage with the latter is that only about 350 bases have to be sequenced compared to 1,500 bases for the 16S rRNA sequence.

#### Table 2: Bacterial strains

	Strain name (name of other collection)	Nucleotides in variable positions	
Bacteria	[accession number of corresponding ssrA	(156, 208 and 235) of the ssrA gene of <i>L. lacti</i> s	
	gene nucleotide sequence]		
Lactococcus lactis ssp.	IL2961 (NCDO2091) [AF375554]	AGT	
lactis	IL2962 (NCDO2110) [AF375555]	AGC	
	IL2965 (NCDO2118) [AF375552]	AGC	
	IL638 [AF375557]	AGC	
	CNRZ337 [AF375561]	AGC	
Lactococcus lactis ssp. lactis	IL890 [AF375558]	AGC	
biovar.diacetylactis	IL986 [AF375562]	TGC	
,	IL1403 <sup>*</sup> [AF375553]	TGC	
	CNRZ124 [AF375566]	TGC	
	CNRZ188 [AF375565]	TGC	
Lactococcus lactis ssp.	IL582 <sup>**</sup> [AF375556]	AAT	
cremoris	IL960** [AF375560]	AAT	
	CNRZ112 [AF375563]	AAT	
	CNRZ119 (ATCC19257) [AF375564]	AAT	
	CNRZ355 [AF375559]	AAT	
Lactococcus plantarum	CNRZ1322 (ATCC43199) [AF375567]		
Lactococcus raffinolactis	CNRZ1214 [AF375578]		
Lactococcus garvieae	CNRZ1323 [AF375579]		
Streptococcus thermophilus	CNRZI358 (ATCC19258) [AF375568]		
Leuconostoc lactis	CIPI02422 (ATCC19256) [AF375569]		
Leuconostoc pseudomesenteroides	CIPI03316 (ATCC12291) [AF375570]		
Leuconostoc mesenteroides	CNRZ361 (ATCC19254) [AF375571]		
ssp. cremoris			
Leuconostoc mesenteroides	CNRZ1022 [AF375572],		
ssp. mesenteroides	CNRZ749 (ATCC8293) [AF375573]		
Leuconostoc mesenteroides	CNRZ77 (ATCC19255) [AF375574]		
ssp. dextranium			
Lactobacillus helveticus	CNRZ223 (ATCC15009) [AF375575]		
Lactobacillus plantarum	CNRZ211 (ATCC14917) [AF375576]		
Lactobacillus gallinarum	CNRZ1931 (ATCC33199, CIP103611) [AF375577]		
Enterococcus faecalis	CNRZI666 (ATCC19433) [AF375580]		
Enterococcus faecium	LMG11423 [AF3755781]		
Enterococcus durans	CNRZ129 (ATCC19432) [AF375582]		
Listeria innocua	CLIP8811 (ATCC33090) [AF375583]		
Listeria monocytogenes	ATCC15313 [AF375584]		
Staphylococcus saprophyticus	ADN12 [AF375585]		
Staphylococcus epidermidis	DSM20044 [AF375586]		
Staphylococcus xylosus	DSM20266 [AF375587]		
Bacillus subtilis	168 [AF375588]		
Escherichia coli K 12	W310.W3110 4 ccr4		

\*Strain IL1403 does not produce diacetyl as it was cured of plasmids [38]. However, the parental strain CNRZ157 (IL594) was identified as biovar. *diacetylactis*, since it produces 7.4 mg/l of diacetyl (Tremblay J, Tailliez P, unpublished). Therefore we consider IL1403 as biovar. *diacetylactis*. \*\* These strains exhibit ssp. *lactis* phenotype (Tremblay J, Tailliez P, unpublished).

For reconstruction of a tmRNA-based phylogenetic tree, we aligned our sequences to other available *ssrA* sequences [16,17]. Depending on the treeing method and whether a filter was used or not, the resulting trees differed in the branching order of particular clusters, although the clusters themselves remained stable (data not shown). This finding is reflected in the multifurcations drawn in the consensus tree (Fig. 2). A stable branching order could not be maintained indicating that the resolution of tmRNA might be restricted to subspecies, species or genus level due to high variability. This was also evident when calculating the 50% conservation filter that left only 180 bases for further analyses. For a thorough phylogenetic study above the level of intraspecies or intrageneric relationships, the tmRNA sequences lack sufficiently high numbers of more conservative positions as one of the prerequisites for phylogenetic markers [24]. However, at the level of genus, species and subspecies they might be an alternative to rRNA sequences that are too conserved for a proper resolution at that level.

The presence of conserved ends on the DNA sequence facilitated the design of universal primers for PCR amplification of the near full-size *ssrA* gene of numerous Grampositive bacteria. The small size of the molecule is an attractive characteristic since it enabled using the same two primers for sequence determination. The use of PCR amplification and sequencing of tmRNA genes was also demonstrated for other groups of bacteria [10,25]. Based on the above analyses we propose that tmRNA sequences can be used for identification of Gram-positive bacteria on the species and genus level. For some bacteria, e.g., lactococci, differences in tmRNA sequences make strain identification possible on the subspecies level, especially if sensitive techniques such as DGGE or TGGE are used [26].

## FISH with the tmRNA targeted probes

*E.* coli W3110 and its  $\Delta ssrA$  derivative were used to test whether it is possible to specifically visualize bacteria by means of FISH with tmRNA targeted probes. Because of relatively low tmRNA molecule numbers per cell (estimated at ~1,000; [11]) HRP-tyramide signal amplification system was applied [27]. We were able to visualize strain W3110 with the tmRNA targeted HRP-labeled probe (Fig. 4, b). The signal was specific, since the  $\Delta ssrA$ strain did not give any signal with the same probe (Fig. 4, d). These results demonstrate that tmRNA, fluorescently labeled after HRP-tyramide signal amplification can be used to distinguish E. coli strains expressing or not the ssrA gene. The ability to visualize bacterial cells also indicates that despite its small size the tmRNA molecule was not totally washed out of the cell during FISH procedure. Using a less sensitive technique with fluorescein labeled probes (E1 - E5, Table 1) we detected a signal only with the strain W3110  $\Delta ssrA$  carrying the ssrA gene on plasmid p10SA, but not with the wild type strain. However, no signal was obtained with wild type strain W3110, even if using all five probes simultaneously.

The low efficiency of fluorescein labeled probes may be due to low copy number or poor accessibility of target RNA. The probes used in our study were targeted to regions with different degree of accessibility [28]. However, using these probes for FISH with the plasmid carrying strain, we did not observe differences in signal intensity (data not shown). It is likely that the tmRNA copy number per cell is too low to be detected by FISH using fluorescein labeled probes, which may restrict its use as a target molecule.

Besides rRNA and regulatory RNAII of ColE1-related plasmids, tmRNAs are the only molecules in bacteria that have been thus far detected by FISH with fluorophore- or HRP-labeled oligonucleotide probes [29]. All three types of molecules are non-translated RNAs. Labeling of specific mRNA with FISH did not give any signal in similar conditions despite strong expression of the selected gene (our unpublished results). mRNA was visualized by FISH only with longer, digoxigenin labeled transcript probes followed by antibody detection [30– 32]. Poor performance of oligonucleotide probes in detection of mRNA may be attributed to involvement of mRNA in the translation process.

## Conclusions

We determined the sequences of tmRNA genes in numerous Gram-positive bacteria. The presence of conservative regions at extremities of tmRNA genes that flank divergent sequences and the possibility to detect tmRNA by FISH makes this molecule a potential tool for identification of bacteria.

## Materials and methods

## Strains, media and general methods

The bacterial strains used in this study are listed in Table 2. E. coli and B. subtilis were grown on L-Broth and other bacteria on BHI broth (Difco, Detroit, Mich.). Solid medium contained 1.5% Bacto-Agar (Difco, Detroit, MI). Standard procedures were used for DNA preparation, restriction endonuclease digestion and DNA electrophoresis [33]. Plasmid p10SA, encoding the ssrA gene of E. *coli*, and strain W3110 $\Delta$ *ssrA* were kindly provided by Dr. H. Inokuchi [12]. B. subtilis strain 168 is from the collection of Dr. C. Anagnostopulos [34]. Chromosomal DNA of staphylococci was obtained from M. C. Montel (France). Other strains are from the CNRZ collection (INRA, Jouy-en-Josas, France). The Lactococcus lactis ssp. lactis strains were considered belonging to biovariety diacetylactis if they produced more that 4 mg/l diacetyl (P. Tailliez and J. Tremblay, unpublished).

## Oligonucleotide probes

The nucleotide sequences of probes are listed in Table 1. These probes were labeled with either fluorescein (F-) or horseradish peroxidase (HRP-). Fluorescent and non-labeled probes were purchased from Eurobio (Les Ulis, France). The probes were labeled with horseradish peroxidase as described before [27].

## DNA sequence analysis

To obtain DNA fragments carrying the *ssrA* gene coding for tmRNA we used PCR amplification. Reactions (94°C



#### 0.

## Figure 2

Consensus tree based on tmRNA sequences determined in this study. For details on the *Lactococcus lactis* group see Table 2. The sequence of *E. coli* was used for setting the root. The bar represents 10% estimated sequence divergence. \*Sequences of bacteria presented without a strain number were taken from the tmRNA databases (references 16, 17).

for 5 min., 50°C for 1 min., 72°C for 7 min., 30 cycles) were performed on Perkin Elmer GeneAmp System 2400 (Perkin Elmer, Norwalk, CT) with Taq DNA polymerase (Promega, Madison, WI). We used primers 11L and 12L for PCR amplification of the internal part of *ssrA* gene of 37 strains of Gram-positive bacteria with low G+C content, of the genera *Lactococcus, Lactobacillus, Streptococcus, Staphylococcus, Leuconostoc, Liste-*

*ria* and *Enterococcus*. To obtain the PCR product of corresponding size for strains *L. gallinarum* CNRZ1931, *L. plantarum* CNRZ211 and *S. xylosus* DSM20266, we had to lower the stringency of PCR (annealing temperature 30°C instead of 50°C), which may indicate that the primers were not fully complementary to target sequences. PCR products were run on agarose gel, the fragment of corresponding size was cut from the gel, purified using



## Figure 3

Epifluorescence micrographs of *E. coli* cells, hybridized with HRP-labeled tmRNA targeted probe (called RNA1) and colored with DAPI; without filter for excitation light (a, c); with a filter for fluorescein (DAPI staining excluded, b, d); W3110 cells (a, b), W3110  $\Delta ssrA$  (c, d).

Gene Clean kit (Bio101, Vista, CA) and sequenced using DNA sequencing kit (PE Applied Biosystems, Warrington, UK) using the same primers. DNA sequences were analyzed with Staden sequence analysis package [35].

For a phylogenetic study, the new tmRNA sequences [16,36]were aligned on the basis of the published alignment [36] with the aid of the ARB software package [ht-tp://www.mikro.biologie.tu-muenchen.de/pub/ARB/] that was used as well for the following analyses. A 50% conservation filter was calculated to exclude the most variable positions [24]. Three different treeing methods,

namely maximum likelihood, maximum parsimony and distance matrix were applied. For the latter method only the part of the sequences common to all was used, since it was sensitive to different sequence lengths. Finally, a consensus tree was drawn based on the maximum likelihood tree [24].

## DNA sequence accession numbers

Sequences of the tmRNA gene fragment that was amplified using primers 11L and 12L, are available in GenBank (see accession numbers in Table 2). The sequence of primers is not included in the sequences.

#### **FISH** techniques

Cell samples were fixed with paraformaldehyde, hybridized with either the fluorescent probes or with horseradish peroxidase (HRP) labeled probes with subsequent fluorophore-tyramide (NEN Life Science Products, Boston, MA) coloration as described [27,37]. 4',6-diamino-2-phenylindole (DAPI, Sigma, St. Louis, MO, 2.5  $\mu$ g/ml) was added directly to the glycerol/PBS mountant to visualize all bacteria (Citifluor Ltd., Canterbury, UK). Photographs were taken using an epifluorescence microscope, equipped with a three-band filter set for emission light (Nikon, Tokyo, Japan), using Sensia 400 film (Fuji, Tokyo, Japan). Slides were scanned and processed with Adobe Illustrator 7.0 program (Edinburgh, Scotland).

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