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RNA 2007 13: 671-681 originally published online March 16, 2007

Access the most recent version at doi:[10.1261/rna.309507](https://doi.org/10.1261/rna.309507)

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In silico screening of archaeal tRNA-encoding genes having multiple introns with bulge-helix-bulge splicing motifs

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ABSTRACT

In archaeal species, several transfer RNA genes have been reported to contain endogenous introns. Although most of the introns are located at anticodon loop regions between nucleotide positions 37 and 38, a number of introns at noncanonical sites and six cases of tRNA genes containing two introns have also been documented. However, these tRNA genes are often missed by tRNAscan-SE, the software most widely used for the annotation of tRNA genes. We previously developed SPLITS, a computational tool to identify tRNA genes containing one intron at a noncanonical position on the basis of its discriminative splicing motif, but the software was limited in the detection of tRNA genes with multiple introns at noncanonical sites. In this study, we initially updated the system as SPLITSX in order to correctly predict known tRNA genes as well as novel ones with multiple introns. By a comprehensive search for tRNA genes in 29 archaeal genomes using SPLITSX, we listed 43 novel candidates that contain introns at noncanonical sites. As a result, 15 contained two introns and three contained three introns within the respective putative tRNA genes. Moreover, the candidates completely complemented all the codons of two archaeal species of uncultured methanogenic archaeon, RC-I and *Thermofilum pendens* Hrk 5, with novel candidates that were not detectable by tRNAscan-SE alone.

Keywords: intron-containing tRNA; bulge-helix-bulge (BHB) splicing motif; archaea; SPLITS; tRNAscan-SE; bioinformatics

INTRODUCTION

In archaeal and eukaryal species, transfer RNA (tRNA)-encoding genes (tDNAs), which constitute one of the major noncoding RNA families, have been reported to contain enzyme-dependent spliceable introns. Although most of the introns of eukaryotic tDNAs are located at unique sites in the anticodon loop between nucleotide positions 37 and 38 (referred to as 37/38, or the canonical position), introns of archaeal tDNAs are also located at other positions (non-canonical positions) (Valenzuela et al. 1978; Daniels et al. 1985). In 2003, Marck and Grosjean identified and summarized the predicted locations, RNA structural topologies, and sizes of all introns located on the tDNAs of 18 archaeal chromosomes (Marck and Grosjean 2003). Of the 136 introns in the total of 800 archaeal tDNAs analyzed, 103 are known to be located at position 37/38, and the remaining 33 are located at 14 other sites on tDNAs, such as the

anticodon stem, amino acid arm, D- and T-loops, and V-arm, and six of the tDNAs were reported to harbor two introns (Wich et al. 1987; Kjems et al. 1989; Smith et al. 1997; She et al. 2001; Fitz-Gibbon et al. 2002; Marck and Grosjean 2003).

All of these tDNAs have a bulge-helix-bulge (BHB) structural consensus, which is also formed in most archaeal pre-rRNA and pre-mRNA introns (Tang et al. 2002; Watanabe et al. 2002; Yoshinari et al. 2006) around their exon-intron boundaries (Kaine et al. 1983; Daniels et al. 1985; Datta et al. 1989; Thompson et al. 1989; Kleman-Leyer et al. 1997). The canonical BHB is a single-hairpin structure that consists of two bulges (B) of 3 nucleotides (nt) separated by a central helix (H) of 4 base pairs (bp) within consensus motif sequences (hBHBh'). For introns at locations other than 37/38, canonical hBHBh' motifs are not always formed, but a simplified HBh' motif consisting of two helices (H and h') and only one bulge can be discerned (Marck and Grosjean 2003). In either case, the splicing sites are always located two bases downstream of the central helix, and the introns are spliced out by the RNA endonucleases. Three classes of tRNA splicing endonucleases have been described and characterized in 19

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Article published online ahead of print. Article and publication date are at <http://www.rnajournal.org/cgi/doi/10.1261/rna.309507>.

archaeal genomes (Tocchini-Valentini et al. 2005): they are a heterotetrameric ($\alpha_2\beta_2$) enzyme in the Crenarchaeota, and homodimeric (α'_2) and homotetrameric (α_4) enzymes in the Euryarchaeota. Most tDNA introns located at non-canonical positions have been found in Crenarchaeal genomes that encode heterotetrameric enzymes, and a few others have also been observed in the Euryarchaeota whose genomes encode homotetrameric enzymes (Smith et al. 1997; Slesarev et al. 2002). However, no tDNAs that have introns in noncanonical positions in Euryarchaeota whose genome encodes homodimeric enzymes have been reported to date.

Over the last decade, most research on the prediction and annotation of tDNAs has utilized tRNAscan-SE software (Lowe and Eddy 1997), especially in genome sequencing projects. tRNAscan-SE combines three different tRNA search methods: tRNAscan 1.3 (Fichant and Burks 1991), the Pavesi search algorithm (Pavesi et al. 1994), and covariance model analysis (Eddy and Durbin 1994), in order to enable fast and highly sensitive prediction of tDNAs without introns or with one intron at the canonical position. Because of this optimization for canonical tDNAs, using a stochastic model learned from mature tRNA structures consisting of cloverleaf structural constraints and consensus sequences, tRNAscan-SE cannot correctly identify >60% of tDNAs with noncanonical introns (Sugahara et al. 2006). To complement tRNAscan-SE for the identification of tDNAs with noncanonical introns, we previously developed the SPLITS toolkit (Sugahara et al. 2006), which predicts and determines introns with BHB motifs within each putative tDNA sequence predicted by the Virtual Footprint (Munch et al. 2005), and removes the introns before passing the sequence to tRNAscan-SE. SPLITS has contributed to the recent genome sequencing project of *Cenarchaeum symbiosum* of the Crenarchaea, by the annotation of all tDNAs whose introns are located at non-canonical sites (Hallam et al. 2006). However, SPLITS was unable to detect multiple intron-containing tDNAs whose introns are harbored within the motif regions of tDNAs corresponding to the target sites of Virtual Footprint screening.

In this study, we analyzed 29 archaeal genomes and predicted novel tDNA candidates with multiple introns, and for this purpose we developed SPLITSX, an enhanced, upgraded version of SPLITS. SPLITSX first predicts non-canonical introns from the whole-genome sequence on the basis of the structural prediction of BHB motifs, and the genome sequences after all possible combinational patterns of intron removal are automatically scanned by tRNAscan-SE. We show that the list of comprehensive archaeal tDNAs predicted by SPLITSX contained all documented tDNAs with noncanonical introns reported in the Archaea. Moreover, with the combination of SPLITSX and tRNAscan-SE, we identified a full set of tRNAs corresponding to all 61 sense codons and one initiator codon in the two

archaeal genomes of uncultured methanogenic archaeon RC-I (RC-I) and *Thermofilum pendens* Hrk 5 (*T. pendens*), where the full set of tRNAs is not detectable by tRNAscan-SE alone. The RC-I genome was further identified to encode the homodimeric type of tRNA-splicing endonuclease, which has previously been suggested not to splice noncanonical introns, and we here present the possibility that the noncanonical introns of tRNAs are actually spliced by the homodimeric enzymes. According to the novel candidates harboring multiple introns, we suggest that many types of tRNA splicing exist in archaeal cells.

RESULTS

Comprehensive screening of novel tDNA candidates in 29 archaeal genomes

For the prediction of novel tDNAs with introns at non-canonical positions by SPLITSX, we analyzed 28 archaeal genomes whose sequences were completely assembled, and the draft assembly genome of *T. pendens*. In total, 74 tDNA candidates that contained introns at noncanonical positions and one tDNA candidate that contained an intron at 37/38 were screened from 11 archaeal genomes (Table 1); 67 out of the total of 75 candidates were predicted from six Crenarchaeal genomes (*Aeropyrum pernix* K1, *Pyrobaculum aerophilum*, *Sulfolobus acidocaldarius*, *Sulfolobus solfataricus*, *Sulfolobus tokodaii*, and *T. pendens*), seven candidates were from four Euryarchaeal genomes (*Methanopyrus kandleri*, *Methanothermobacter thermautotrophicus*, *Methanospirillum hungatei*, and RC-I), and one candidate was from the Nanoarchaea, *Nanoarchaeum equitans*. Except for two cases, all novel candidates scored covariance model (COVE) scores (Eddy and Durbin 1994) of >55.00 bits, which was the lowest score of all the documented tDNAs used as positive controls. Furthermore, almost all novel introns were observed to have well-conserved BHB structures whose free energies and position weight matrix (PWM) scores were on a par with those of documented BHB motifs (for details, see Supplemental Materials, Fig. S1, at http://splits.iab.keio.ac.jp/RNA_SupplMat.pdf). The 75 candidates included 32 documented tDNAs, and the remaining 43 were novel candidates. The 32 candidates covered all of the previously documented tDNAs with one or two introns at noncanonical positions. The 43 novel candidates were predicted in the *P. aerophilum*, *S. acidocaldarius*, *T. pendens*, *M. hungatei*, and RC-I genomes. Twenty-four out of the 43 candidates contained a single intron at noncanonical sites, 15 contained two introns, and three contained three introns within a single tDNA gene. Furthermore, 33 out of 43 overlapped with documented tDNA regions predicted by tRNAscan-SE, and these candidates reassigned the locations of introns or anticodons of the tRNAs. Three candidates from the genome sequence

TABLE 1. All 75 tDNA candidates containing noncanonical introns predicted by SPLITSX

Genome sequence	Accession No. ^a	Enzyme ^b	Gene ID ^c	Left boundary ^d	Right boundary ^e	Strand	Isotype	Anticodon	Intron location ^f	COVE
<i>A. pernix</i>	NC_000854	$\alpha_2\beta_2$	AP01*	150239	150360	1	Pro	CGG	32/33	84.43
			AP02*	236316	236428	1	Lys	CTT	45/46	93.13
			AP03*	295231	295325	1	Thr	TGT	22/23	90.42
			AP04*	402990	403103	1	Trp	CCA	30/31	88
			AP05*	1176266	1176378	0	Lys	TTT	45/46	91.48
<i>P. aerophilum</i>	NC_003364	$\alpha_2\beta_2$	PA01*	21776	21867	1	His	GTG	44/45	70.91
			PA02	486449	486542	1	Arg	GCG	56/57	82.91
			PA03	487826	487945	0	Glu	TTC	3/4, 58/59	85.05
			PA04*	663679	663772	0	Ile	CAT	29/30	92.75
			PA05*	837670	837785	1	Tyr	GTA	39/40	78.96
			PA06*	895379	895474	1	Glu	CTC	3/4	85.43
			PA07*	1218974	1219067	0	Met	CAT	29/30	86.6
			PA08*	1226004	1226102	1	Asp	GTC	3/4	81.03
			PA09	1239015	1239132	0	Thr	TGT	29/30, 59/60	91.28
			PA10*	1298180	1298281	1	Arg	CCT	30/31	85.39
			PA11*	1661495	1661585	0	iMet	CAT	38/39	85.48
			PA12	1792081	1792192	1	Thr	GGT	38/39, 56/57	89.74
			PA13*	1797828	1797917	1	Gln	CTG	21/22	74.52
			PA14*	1819184	1819275	1	Val	GAC	29/30	85.5
PA15*	1924495	1924596	0	Arg	CCG	30/31	87.46			
PA16*	2007525	2007609	0	Ser	GGA	49/50	74.69			
PA17*	2028602	2028700	0	Cys	GCA	58/59	87.85			
PA18*	2039014	2039108	0	Thr	CGT	29/30	89.11			
<i>T. pendens</i> (Contig10)	NZ_AASJ01000002	$\alpha_2\beta_2$	TP01*	31543	31646	0	Leu	CAA	30/31	77.99
			TP02	55801	55908	0	iMet	CAT	37/38, 59/60	41.98
			TP03	56190	56310	0	Glu	TTC	20/21, 45/46	83.78
			TP04*	58492	58608	1	Ser	CGA	32/33, 37/38	75.57
			TP05*	58759	58861	1	His	GTG	45/46	81.42
			TP06*	63410	63552	0	Pro	GCG	32/33, 37/38, 45/46	83.34
			TP07*	64151	64249	0	Arg	CCG	45/46	80.7
			TP08*	66393	66489	0	Gly	TCC	45/46	86.17
			TP09*	105499	105620	0	Ile	CAT	22/23, 43/44	85.81
			TP10*	117109	117205	1	Arg	TCT	32/33	89.78
			TP11*	132255	132384	1	Val	TAC	25/26, 37/38	86.72
			TP12*	156314	156411	0	Lys	CTT	30/31	89.49
			TP13	168930	169057	0	Leu	CAG	30/31, 37/38	74.83
			TP14*	170165	170273	1	Phe	GAA	44/45	78.43
			TP15*	174055	174185	1	Cys	GCA	30/31	69.82
			TP16*	200615	200715	1	Arg	TCG	45/46	83.52
			TP17	202868	202990	0	Ala	TGC	25/26, 37/38	79.72
			TP18*	234235	234329	1	Gly	CCC	45/46	88.82
			TP19	273269	273378	1	Leu	TAA	30/31	77.33
			TP20*	274272	274401	1	Asn	GTT	22/23, 43/44	81.66
			TP21*	326646	326754	1	Asp	GTC	24/25, 45/46	79.23
			TP22*	326914	327043	0	Ile	GAT	22/23, 43/44	84.26
			TP23*	327131	327242	1	Trp	CCA	30/31, 45/46	85.43
			TP24*	337454	337557	0	Lys	TTT	30/31	92.86
			TP25*	337638	337762	0	Val	CAC	25/26, 37/38	81.06
			TP26*	352082	352175	0	Val	GAC	28/29	84.8
			TP27*	363819	363921	0	Ser	TGA	32/33	72.61
<i>T. pendens</i> (Contig11)	NZ_AASJ01000001	$\alpha_2\beta_2$	TP28*	5155	5251	0	Arg	CCT	32/33	85
			TP29*	17245	17387	0	Pro	TGG	32/33, 37/38, 45/46	83.9
			TP30*	48570	48705	0	Pro	GGG	25/26, 37/38, 43/44	88.07
			TP31*	56976	57071	1	Arg	GCG	45/46	79.27
			TP32	114663	114787	1	Glu	CTC	20/21, 45/46	85.92
			TP33	169396	169517	0	Ala	CGC	25/26, 40/41	79.89

(continued)

TABLE 1. Continued

Genome sequence	Accession No. ^a	Enzyme ^b	Gene ID ^c	Left boundary ^d	Right boundary ^e	Strand	Isotype	Anticodon	Intron location ^f	COVE
<i>S. acidocaldarius</i>	NC_007181	$\alpha_2\beta_2$	TP34*	174843	174950	1	Ala	GGC	43/44	74.83
			TP35*	201017	201113	0	Gly	GCC	45/46	85.54
			SA01*	395126	395216	0	Glu	TTC	20/21	83.05
<i>S. solfataricus</i>	NC_002754	$\alpha_2\beta_2$	SA02*	694093	694183	0	Glu	CTC	20/21	83.74
			<u>SS01*</u>	142457	142570	1	Cys	GCA	28/29, 37/38	55.39
			<u>SS02*</u>	453232	453321	1	Glu	TTC	20/21	87.41
<i>S. tokodaii</i>	NC_003106	$\alpha_2\beta_2$	<u>SS03*</u>	648224	648313	1	Glu	CTC	20/21	86.85
			<u>ST01*</u>	191161	191251	1	Glu	TTC	20/21	83.05
			<u>ST02*</u>	280511	280608	1	iMet	CAT	38/39	79.12
			<u>ST03*</u>	371083	371173	1	Glu	CTC	20/21	83.22
<i>N. equitans</i>	NC_005213	$\alpha_2\beta_2$	<u>ST04</u>	482554	482657	1	Leu	GAG	30/31	65.14
			<u>NE01</u>	151992	152078	0	Trp	CCA	30/31	85.27
<i>M. kandleri</i>	NC_003551	$\alpha_2\beta_2$	<u>MK01</u>	1413744	1413873	0	Glu	TTC	20/21, 37/38	80.37
			<u>MK02*</u>	1659600	1659729	0	Glu	TTC	20/21, 37/38	81.88
<i>M. thermotrophicus</i>	NC_000916	α_4	<u>MT01*</u>	21281	21403	0	Pro	GGG	32/33, 37/38	54.99
<i>M. hungatei</i>	NC_007796	α'_2	MH01*	2881749	2881854	1	His	GTG	34/35, 37/38	71.3
RC-I	AM114193	α'_2	UM01	358273	358368	1	Ile	GAT	23/24	80.23
			UM02	358484	358579	1	Ile	GAT	23/24	80.23
			UM03	2964519	2964767	0	Trp	CCA	37/38	71.52

Candidates for tRNA-encoding genes including noncanonical introns in archaeal species are listed along with their enzyme architecture, position, strand, amino acid charge, anticodon, location of introns, and COVE score calculated by tRNAscan-SE.

^aDenotes the GenBank accession number of the target species.

^bThe documented or predicted enzyme architecture of the tRNA endonucleases.

^cDocumented tRNA-encoding genes are indicated by underlining, and overlapping candidates are indicated by asterisks.

^dThe 5'-end position on the plus strand "0" or the 3'-end position on the minus strand "1."

^eThe 3'-end position on the plus strand "0" or the 5'-end position on the minus strand "1."

^fMultiple introns of the candidates are divided by commas.

of *S. acidocaldarius* (SA01 and SA02) and *P. aerophilum* (PA01) were in agreement with our previous work (Sugahara et al. 2006).

Predicted set of tRNA genes fulfills all 61 codons in *T. pendens*

By the analysis of the genome sequence of *T. pendens* by tRNAscan-SE, a total of 39 tDNAs were predicted (Table 2). The candidates predicted by tRNAscan-SE alone missed a number of tDNAs corresponding to 15 sense codons and one initiator Met tDNA (tDNA-iMet) (Fig. 1A). On the other hand, SPLITSX revealed a total of 35 tDNA genes that contained single or multiple introns at noncanonical sites. Seven out of 35 were predicted from novel genomic regions, and all of the remaining 28 candidates overlapped with tDNA regions predicted by tRNAscan-SE with higher COVE scores. Moreover, the annotations of the corresponding amino acids of 20 tDNAs predicted by tRNAscan-SE alone were reassigned with reliable COVE scores. Combining the 11 candidates predicted only by tRNAscan-SE and the novel 35 candidates by SPLITSX, we identified a total of 46 functional tDNA candidates that were able to read 59 sense codons and one initiator codon AUG, considering the hybridization of dG and rU at a wobble position.

However, the candidates generated by the merger of the results of SPLITSX and tRNAscan-SE missed only one tDNA encoding tRNA^{Ser} (GCU), which reads sense codons of AGC and AGU. Therefore, we further analyzed the *T. pendens* genome with SPLITSX with more relaxed parameters, and the putative tRNA^{Ser} (TP36) encoded at a genomic location between 200101 and 200210 of Contig 10 (NZ_AASJ01000002) on the complementary strand was identified. The anticodon of TP36 was GCU, the COVE score was 50.35, and the intron was located at 21/22 (for detailed structure, see Supplemental Fig. S1 at http://splits.iab.keio.ac.jp/RNA_SupplMat.pdf). TP36 overlapped with a tRNA^{Ala}-encoding gene (SE17) containing an intron located at 37/38 that was predicted by tRNAscan-SE with a COVE score of 64.18. However, the intron (37/38) of SE17 predicted by tRNAscan-SE could not form any type of BHB motifs. In contrast, the 20-nt intron of TP36 predicted by SPLITSX formed a canonical hBHBh' motif with a stable free energy of -3.20 kcal/mol and high PWM scores of 0.83, and was located at position 21/22, which is reported to harbor a 17-nt intron. Therefore, we suggest that TP36 actually exists in the *T. pendens* cells to encode tRNA^{Ser} (GCU), rather than as SE17 encoding tRNA^{Ala}. Accordingly, the above-mentioned set of tDNAs fulfills all 61 sense codons and the initiator codon AUG (Fig. 1B).

TABLE 2. The tDNA candidates in *T. pendens* predicted by tRNAscan-SE alone and by SPLITSX

Accession No.	tRNAscan-SE						SPLITSX					
	Gene ID	Genome loci	Strand	Isotype	Anticodon	COVE	Gene ID	Genome loci	Strand	Isotype	Anticodon	COVE
NZ_AASJ01000003	SE01+	188076–188152	1	Gln	TTG	79.61						
NZ_AASJ01000002	SE02	31568–31646	0	PSE	CAA	22.85	TP01+	31543–31646	0	Leu*	CAA	77.99
	SE03+	56010–56097	1	Ser	GGA	73.43	TP02+	55801–55908	0	iMet	CAT	41.98
	SE04	58492–58608	1	Lys	CTT	58.99	TP03+	56190–56310	0	Glu	TTC	83.78
	SE05	58759–58861	1	His	GTG	60.27	TP04+	58492–58608	1	Ser*	CGA*	75.57
	SE06	63410–63552	0	Ser	CGA	61.97	TP05+	58759–58861	1	His	GTG	81.42
	SE07	64151–64249	0	Arg	CCG	65.53	TP06+	63410–63552	0	Pro*	CGG*	83.34
	SE08	66393–66489	0	Gly	TCC	75.54	TP07+	64151–64249	0	Arg	CCG	80.7
	SE09	105499–105620	0	PSE	TAC	38.73	TP08+	66393–66489	0	Gly	TCC	86.17
	SE10	117109–117205	1	Ala	CGC	69.37	TP09+	105499–105620	0	Ile*	CAT*	85.81
	SE11+	129907–129994	1	Leu	GAG	74.75	TP10+	117109–117205	1	Arg*	TCT*	89.78
	SE12	132255–132384	1	PSE	CAC	45.33	TP11+	132255–132384	1	Val*	TAC*	86.72
	SE13+	134467–134543	1	Gln	CTG	75.58						
	SE14	156314–156411	0	UND	???	66.56	TP12+	156314–156411	0	Lys*	CTT*	89.49
							TP13+	168930–169057	0	Leu	CAG	74.83
	SE15	170165–170273	1	Phe	GAA	63.43	TP14+	170165–170273	1	Phe	GAA	78.43
	SE16	174055–174185	1	Trp	CCA	47.83	TP15+	174055–174185	1	Cys*	GCA*	69.82
	SE17+	200101–200210	1	Ala	CGC	64.18						
	SE18	200615–200715	1	Arg	TCG	61.6	TP16+	200615–200715	1	Arg	TCG	83.52
							TP17+	202868–202990	0	Al	TGC	79.72
	SE19+	233238–233316	0	Met	CAT	87.15						
	SE20	234235–234329	1	Gly	CCC	73.84	TP18+	234235–234329	1	Gly	CCC	88.82
							TP19+	273269–273378	1	Leu	TAA	77.33
	SE21	274272–274401	1	Leu	TAG	48.99	TP20+	274272–274401	1	Asn*	GTT*	81.66
	SE22	326646–326754	1	PSE	GGA	40.5	TP21+	326646–326754	1	Asp*	GTC*	79.23
	SE23	326914–327043	0	PSE	TAG	51.66	TP22+	326914–327043	0	Ile*	GAT*	84.26
	SE24	327131–327242	1	UND	???	53.13	TP23+	327131–327242	1	Trp*	CCA*	85.43
	SE25+	337204–337304	0	Thr	TGT	77.32						
	SE26	337454–337557	0	Pro	GGG	71.09	TP24+	337454–337557	0	Lys*	TTT*	92.86
	SE27	337638–337762	0	PSE	GGC	52.68	TP25+	337638–337762	0	Val*	CAC*	81.06
	SE28	352082–352175	1	Thr	TGT	63.62	TP26+	352082–352175	1	Val*	GAC*	84.8
	SE29	363819–363921	0	Glu	CTC	58.92	TP27+	363819–363921	0	Ser*	TGA*	72.61
NZ_AASJ01000001	SE30	5155–5251	0	Ala	CGC	65.29	TP28+	5155–5251	0	Arg*	CCT*	85
	SE31	17245–17387	0	Ser	CGA	61.62	TP29+	17245–17387	0	Pro*	TGG*	83.9
	SE32	48570–48705	0	PSE	AAA	50.78	TP30+	48570–48705	0	Pro*	GGG*	88.07
	SE33+	48799–48886	0	Leu	TAG	71.44						
	SE34	56976–57071	1	Arg	GCG	65.24	TP31+	56976–57071	1	Arg	GCG	79.27
							TP32+	114663–114787	1	Glu	CTC	85.92
	SE35+	147962–148062	0	Thr	CGT	77.4						
							TP33+	169396–169517	0	Ala	CGC	79.89
	SE36	174843–174950	1	Ala	GGC	61.26	TP34+	174843–174950	1	Ala	GGC	74.83
	SE37	201017–201113	0	Gly	GCC	74.9	TP35+	201017–201113	0	Gly	GCC	85.54
	SE38+	246716–246808	0	Thr	GGT	67.55						
	SE39+	297647–297778	0	Tyr	GTA	28.25						

Candidates for tRNA-encoding genes in *T. pendens* predicted by tRNAscan-SE alone and by SPLITSX are listed along with their gene ID, genomic loci, strand, isotype of amino acid charge, anticodon, and COVE scores calculated by tRNAscan-SE, respectively. A candidate whose genomic position overlapped with a candidate predicted by another software was shown in same row, and the "Gene ID" of an overlapping candidate whose COVE score was higher than another was checked by "+." For details, see the legend of Table 1.

Structures and isotypes of predicted tDNAs

The locations and lengths of the introns, the structures of their BHB motifs, and the isotypes of the predicted tDNAs are summarized in Figure 2. SPLITSX predicted 54 novel

introns that were located at noncanonical sites of tDNAs, and ~65% (36 of 54) of the introns were located at previously reported tDNA sites, whereas the remaining ~35% (18 of 54) of introns were distributed at eight unreported sites, such as in nucleotide positions 23/24, 24/25, 25/26,

A

		2nd base								
		A	G	C	U					
1st base	A	Lys	0	Arg	0	Thr	2	Ile	0	A
			1				1	Met	1	G
		Asn	0	Ser	0	Thr	1	Ile	0	C
	G	Glu	0	Gly	1	Ala	0	Val	0	A
			1		1		3			G
		Asp	0	Gly	1	Ala	1	Val	0	C
	C	Gln	1	Arg	1	Pro	0	Leu	2	A
			1		1					G
		His	1	Arg	1	Pro	1	Leu	1	C
	U	STOP		STOP		Ser	0	Leu	0	A
				Trp	1		2			G
		Tyr	1	Cys	0	Ser	1	Phe	1	C
									U	

B

		2nd base								
		A	G	C	U					
1st base	A	Lys	1	Arg	1	Thr	1	Ile	1	A
			1		1		1	Met	1	G
		Asn	1	Ser	1	Thr	1	Ile	1	C
	G	Glu	1	Gly	1	Ala	1	Val	1	A
			1		1		1		1	G
		Asp	1	Gly	1	Ala	1	Val	1	C
	C	Gln	1	Arg	1	Pro	1	Leu	1	A
			1		1		1		1	G
		His	1	Arg	1	Pro	1	Leu	1	C
	U	STOP		STOP		Ser	1	Leu	1	A
				Trp	1		1		1	G
		Tyr	1	Cys	1	Ser	1	Phe	1	C
									U	

FIGURE 1. Codon tables of *T. pendens* showing the numbers of tRNAs of respective codons. The *leftmost* column, the *top* row, and the *rightmost* column indicate the first, second, and third bases of sense codons, and their coding amino acids are shown in respective fields. (A) Codon table including the 39 tDNA candidates predicted by tRNAscan-SE alone. (B) Codon table including the 46 tDNA candidates complemented and reassigned by SPLITXS, with 10 candidates identified only by tRNAscan-SE.

34/35, 40/41, 43/44, 44/45, and 49/50. The schematic diagrams of three typical tDNAs that contain introns at unreported sites are displayed in Figure 3A–C. The gene encoding tRNA^{Asp} (GUC) in *T. pendens* had two introns at noncanonical positions, 24/25 and 45/46 (Fig. 3A). The intron located at the unreported region 24/25 formed a typical hBHBh' motif, whereas the other intron located at 45/46 formed an HBh' motif. The gene encoding tRNA^{Ile} (with anticodon CAU reading AUA) also had two introns at 22/23 and 43/44, and both introns formed HBh' motifs (Fig. 3B). In addition, both of those candidates were significant tDNAs that complement gaps of missing tRNAs in *T. pendens*. The gene encoding tRNA^{His} in *M. hungatei* had a 17-nt intron at the unreported site of 34/35, forming an HBh' structure, and a 15-nt intron at the canonical site of 37/38, forming an hBHBh' (Fig. 3C). The intron at 37/38 was found to form the hBHBh' motif after removal of the intron at 34/35. Moreover, UM01 and UM02 in RC-I were analogous genes encoding tRNA^{Ile} (GAU); both had 21-nt introns at unreported sites of 23/24, and UM03 encoding tRNA^{Trp} (CCA) harbored the longest 175-nt intron at 37/38. All of these species of tDNA candidates had not been previously annotated by tRNAscan-SE, and our candidates fulfill all codons in RC-I.

Surprisingly, every proline-charged tDNA of TP06, TP29, and TP30 in the *T. pendens* genome harbored three introns. TP06 and TP29 were analogous tRNA^{Pro} genes that presumably contained three endogenous introns located at 32/33, 37/38, and 45/46. The intron located at the noncanonical site of 32/33 formed an hBHBh' motif, and another noncanonical intron located at 45/46 formed an HBh' motif. The intron located at canonical position 37/38 was found to form an hBHBh' motif after the removal of

the outer two noncanonical introns. A schematic representation of the predicted synthetic procedure of the TP29 pre-tRNA is displayed in Figure 4.

Identification of tRNA-splicing endonuclease subunits

In order to discuss the novel candidates having non-canonical introns from the viewpoint of the types (homodimer, homotetramer, or heterotetramer) of tRNA-splicing endonuclease subunits, we identified tRNA-splicing endonuclease subunits by using BLASTP (Altschul et al. 1990) searches from the respective genomes. Each type of subunit has been described in the Introduction. The types of tRNA-splicing endonuclease subunits and related genes in seven of the listed 11 genomes have already been identified by Tocchini-Valentini et al. (2005), whereas the types of the subunits in the remaining four genomes (*T. pendens*, *S. acidocaldarius*, *M. hungatei*, and RC-I) have not been identified. We therefore additionally searched for genes homologous to those encoding the tRNA-splicing endonuclease subunits of *Methanocaldococcus jannaschii*. A set of two homologs was revealed in each of the genomes of *T. pendens* and *S. acidocaldarius*. In *T. pendens*, one was observed in the genomic region between positions 18524 and 19153 in the direct strand of genomic contig 10, with an expectation value (E-value) of 1.3e-24, and the other was in the region between positions 316950 and 317531 in the direct strand, with an E-value of 1.6e-16. On the genome sequence of *S. acidocaldarius*, one was conserved in the region between positions 687428 and 687973 of the complementary strand, with an E-value of 2.4e-19, and the other was in the region between positions 526525 and 526800 of the complementary strand, with an E-value of

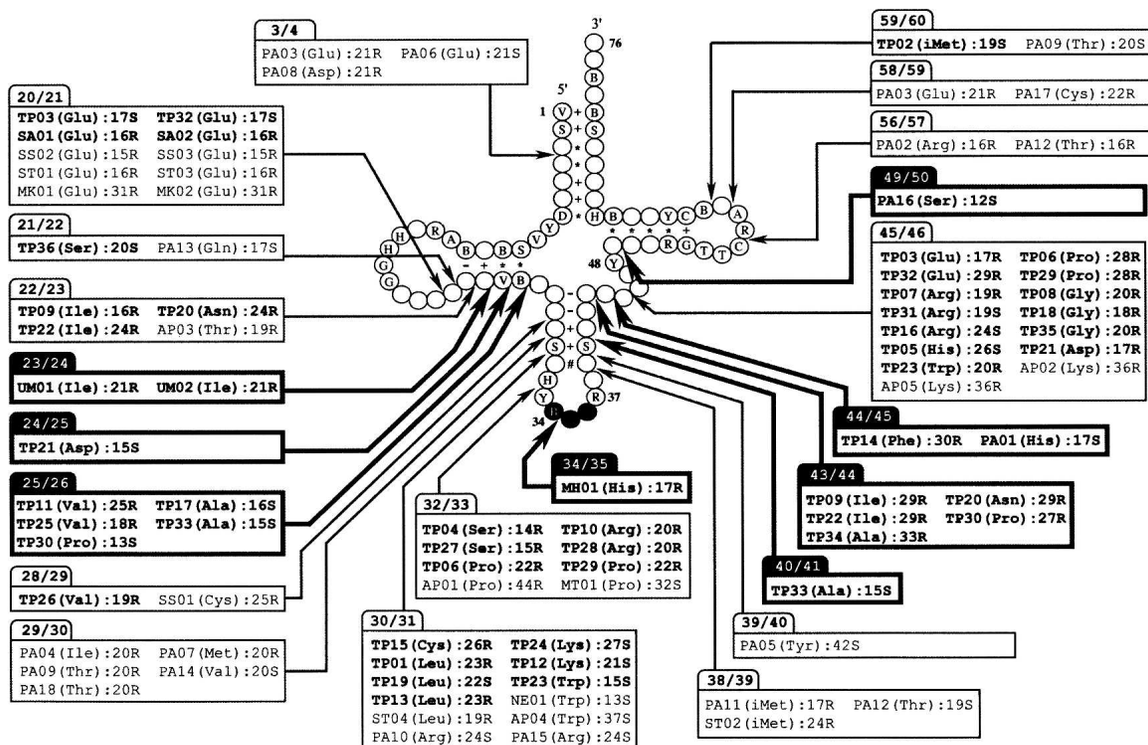


FIGURE 2. Locations of introns of predicted tDNA genes in 11 Archaea. The schematic cloverleaf structure corresponding to the consensus sequence of archaeobacterial tRNA sequences has been modified in accordance with the results of a previous work (Marck and Grosjean 2003). The conventional IUB/UPAC degenerate DNA alphabet is used in this figure: R (purine), A or G; Y (pyrimidine), C or T; S (strong), G or C; B (not A), C, G, or T; D (not C), A, G, or T; H (not G), A, C or T; V (not T), A, C, or G; N (any), A, C, G, or T. Base-pairing consensus is denoted by: (+) Watson-Crick base pairing only; (*) Watson-Crick or G-T/T-G pairings; (#) Watson-Crick pairing or mismatch; (-), Watson-Crick pairing or G-T/T-G pairings or mismatches. Intron positions of documented or novel candidates are shown by thin and bold arrows, and by clear and solid tabs over the text boxes, respectively. Each candidate is listed in the text boxes along with the isotype of the amino acid charge in parentheses. The intron length and type of bulge-helix-bulge (BHB) structure are indicated to the right of the colons. The number denotes the nucleotide length, and the capital letter denotes the type of BHB structure: (S) strict hBHB' motif, (R) relaxed HBh' motif. Novel candidates are indicated by bold text.

7.5e-05. This suggested that the enzyme structures of the tRNA endonuclease in *T. pendens* and *S. acidocaldarius* were heterotetrameric ($\alpha_2\beta_2$). On the other hand, two closely located homologous sequences of lengths of ~150–200 residues were observed in the RC-I genomic DNA between positions 317008 and 318069 of the direct strand with E-values 6.8e-25 and 4.6e-21, and in *M. hungatei* genomic DNA between positions 1954717 and 1955724 of the complementary strand with E-values 6.1e-17 and 5.3e-13, which possibly function as one gene, translating to be the homodimer (α'_2) with ~350 amino acid residues. In the results of multiple alignments with documented endonuclease sequences, the novel sequences have been observed with consensus sequences and strong similarities (see Supplemental Fig. S2 at http://splits.iab.keio.ac.jp/RNA_SupplMat.pdf). Except for UM01 and UM02 of RC-I and MH01 of *M. hungatei*, whose host genomes were predicted to encode homodimeric enzymes, all of the novel tDNA candidates were found in the genomes encoding heterotetrameric or homotetrameric enzymes, which have been

reported to recognize introns located at noncanonical positions in tDNAs. We summarize each type of subunit in Table 1.

DISCUSSION

We have described the comprehensive screening of intron-containing tDNAs, including all of the documented tDNAs harboring noncanonical introns with a reasonable number of candidates within the archaeal species analyzed in this work, using enhanced software designated SPLITSX to detect tRNA-encoding genes with multiple introns. Although SPLITSX functions by simply removing BHB motifs from the genome sequence before the execution of tRNAscan-SE, in the absence of any information on the tRNA sequences or their structural specifications and their genomic loci, ~65% of the predicted introns were located at the sites of previously documented tDNAs. Moreover, ~74% (32 of 43) of novel tDNA candidates were predicted with their analogous candidates that encode tRNAs of the

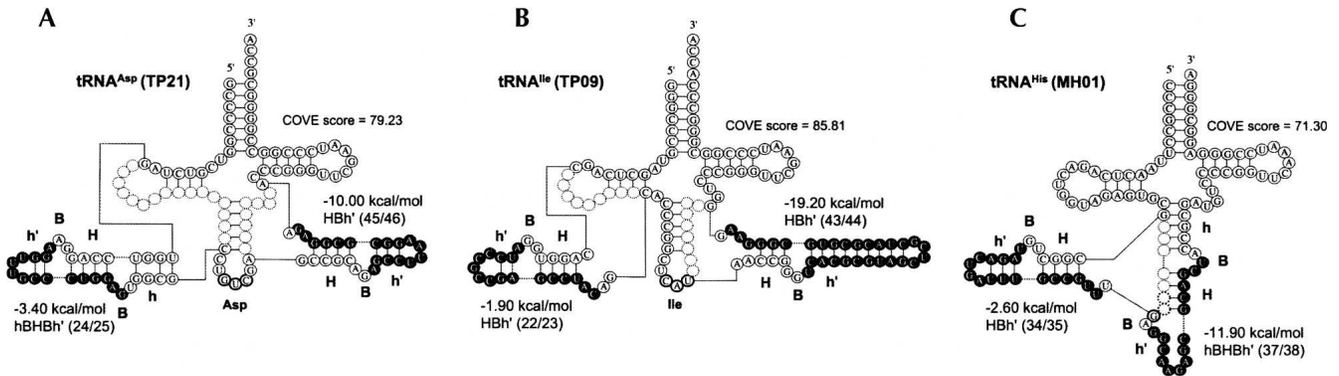


FIGURE 3. Schematic diagrams of novel tRNA-encoding genes including multiple introns predicted by SPLITSX, accompanied by COVE scores and free energies of BHB motif. Intron and exon sequences are represented by solid and clear circles, respectively. (Arrows) Location where introns are located. (A) TP21, a putative tRNA^{Asp} with two introns located between nucleotide positions 24 and 25 (24/25) and 45/46 encoded in the *T. pendens* genomic DNA. The respective position weight matrix (PWM) scores of the BHB motifs were 0.86 and 0.88. (B) TP09, a tRNA^{Ile} with two introns at 22/23 and 43/44 synthesized according to the BHB motif with PWM scores 0.85 and 0.63, in *T. pendens*. (C) MH01, tRNA^{His} with double introns at 34/35 and 37/38 synthesized according to the BHB motif with PWM scores of 0.67 and 0.90, in the *Methanospirillum hungatei*.

same isotypes with similar intron lengths and locations. For example, in addition to the known tDNAs that charge glutamic acids containing introns of 15–16-nt conformations located at 20/21, the novel candidates SA01, SA03, TP03, and TP32 were also suggested to charge glutamic acids, containing introns at 20/21 with lengths of 16–17 nt. Additionally, some candidates that presumably contain introns at undocumented sites were also identified with their paralogous or orthologous genes. For example, UM01 and UM02 both encoded tRNA^{Ile} (GAU), whose introns were 21-nt long and formed HBh' structures located at 23/24. According to the results, we therefore suggest that SPLITSX is able to screen reliable tDNA candidates containing introns. Moreover, SPLITSX also detected UM03,

which contains a 175-nt intron at canonical position 37/38, from the RC-I genome; this is longer than the 121-nt intron previously reported in the *Aeropyrum pernix* genome as the longest intron in archaeal tDNAs.

Our novel 43 candidates reassigned 33 tRNA genes, which had been previously documented mainly by utilizing tRNAscan-SE. Here, we suggest that the reassigned tDNAs are more reliable than the previously annotated ones. For example, a tRNA^{His}-encoding gene had been annotated to contain an intron at 37/38 by Marck and Grosjean (2003), whereas SPLITSX reassigned the tRNA^{His}-encoding gene as PA01, also encoding tRNA^{His} in the same genomic locus, containing an intron at the site of 44/45. Although the previous tRNA^{His} contained a relaxed hBH motif and was

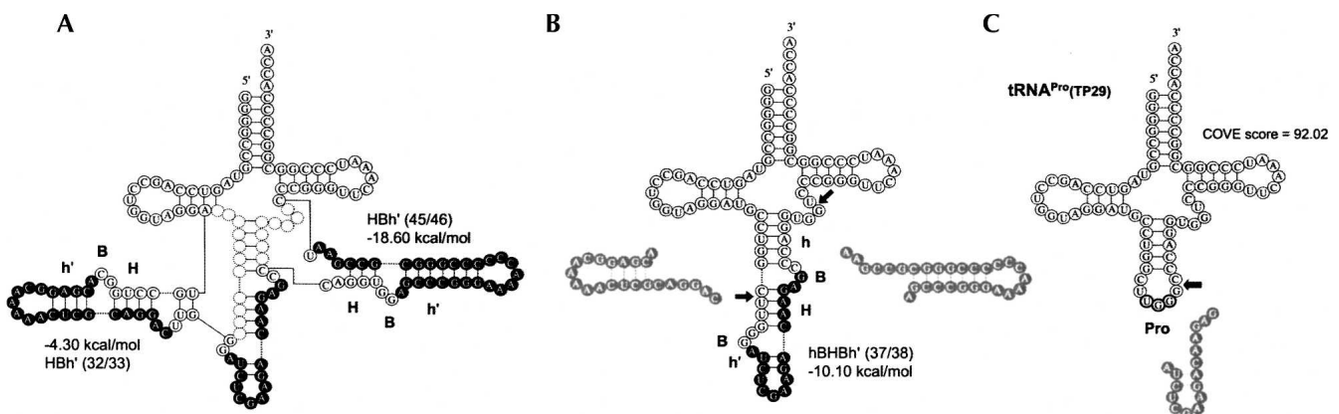


FIGURE 4. Schematic representation of the synthesis procedure in the maturation of tRNA candidate TP29, with three introns located at 32/33, 37/38, and 45/46, along with the COVE score and the free energies of the BHB motifs. (Arrows) Locations where introns were inserted. The position weight matrix (PWM) scores of the respective BHB motifs were 0.74, 0.87, and 0.88. Intron and exon sequences are represented by solid and clear circles, respectively. Two introns that have the potential to form BHB motifs might initially be spliced out at positions 32/33 and 45/46 (A), then another BHB motif is predicted to occur and to be spliced out at position 37/38 (B). Finally, the putative tDNA region of TP29 may form tRNA^{Pro} (C).

located at 37/38, the first helix (h) had one mismatch and the central 4-bp H helix had many rG and rU hybridizations (UUGU/GCGG). On the other hand, the intron located at 44/45 of PA01 predicted by SPLITSX forms a strict hBHBh' motif with no mismatch, and the central H helix is more convincing (CCCG/CGGG instead of UUGU/GCGG). Moreover, the cloverleaf structure of the reassigned candidate is more mature, with a general tRNA structure consisting of a 5-bp stem at the anticodon arm and a 7-nt anticodon loop. The previous tRNA^{His} contained the intron at 37/38, and its mature tRNA consisted of a 4-bp anticodon stem and an oversized 9-nt anticodon loop. Therefore, the COVE score was also lower in the previous structure compared with our reassigned candidate (Fig. 5). We thus suggest that PA01 (tRNA^{His}) containing the intron at 44/45 is actually transcribed, rather than the previous

tRNA^{His}-encoding gene containing the intron at 37/38. Similarly, we claim that other tDNA candidates screened by SPLITSX are more convincing than the previously annotated tDNAs.

We computationally screened a total of 17 tDNAs with multiple introns, including three tDNAs with three introns, encoded in the *T. pendens* genome. However, only six tDNAs throughout all the archaeal species had been reported to contain two introns within a single gene (Marck and Grosjean 2003), and thus *T. pendens* is suggested to preferentially encode many tRNA genes with multiple introns compared with other Archaea. In the process of maturation of the pre-tRNA^{Pro} of *M. thermautotrophicus*—the first documented case containing two introns within a single gene (included in our list as MT01)—the first intron, with an hBHBh' motif of 32 nt located at positions 32/33, is

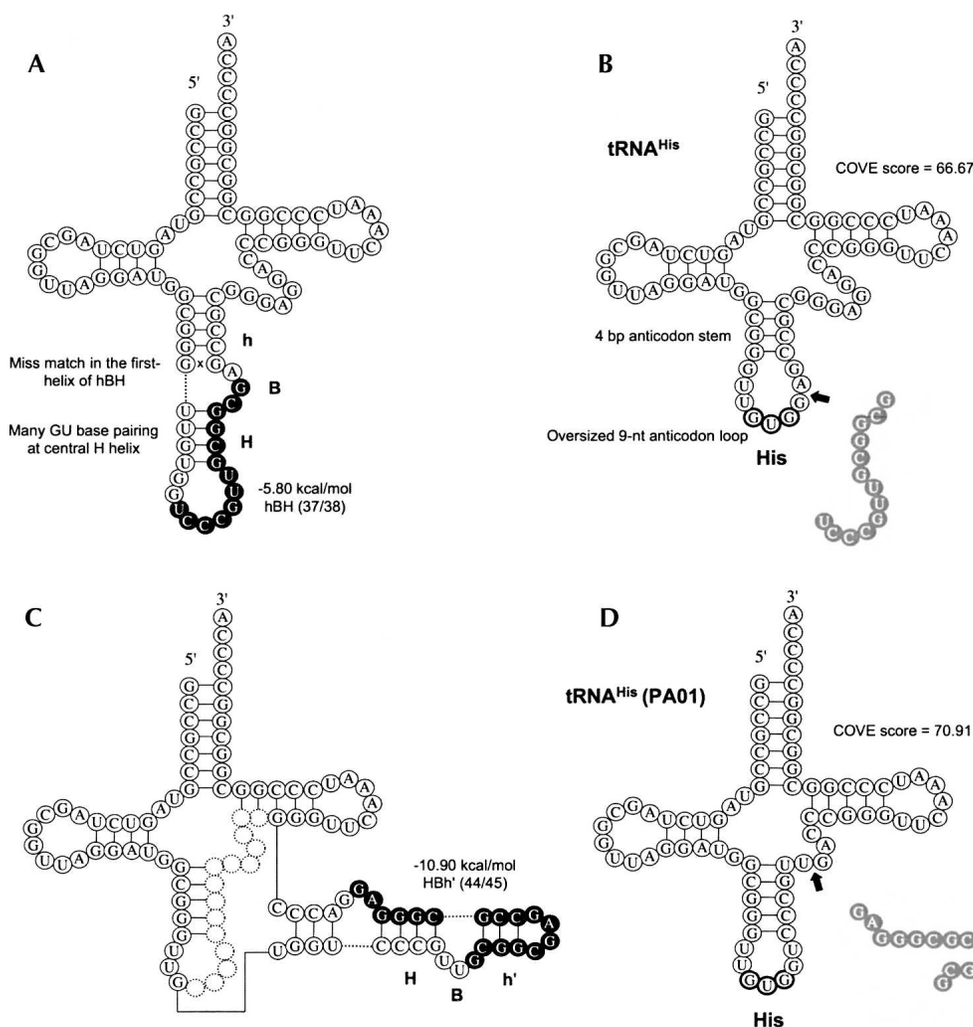


FIGURE 5. Comparison of the *P. aerophilum* tRNA^{His} (GUG) structure previously annotated by Marck and Grosjean (2003) with that reassigned by SPLITSX, along with their COVE scores and free energies of the BHB motif. Intron and exon sequences are represented by solid and clear circles, respectively. (A) The previous pre-tRNA^{His} (GUG) harboring the intron at 37/38 predicted by tRNAscan-SE, (C) the reassigned pre-tRNA^{His} (GUG) harboring the intron at 44/45 predicted by SPLITSX. Mature tRNA structures are displayed in B and D, respectively. (Arrows) Locations where introns were inserted, (x) a mismatch of base pairing in the BHB motif.

initially spliced out, and then the second intron, with an hBHBh' motif of 16 nt, is revealed and processed at the canonical location, position 37/38 (Smith et al. 1997). We speculate that the novel candidate tRNA^{Pro} in *T. pendens* (TP06, TP29, and TP30), with the third intron at position 43/44 or 45/46, is spliced in a similar manner (for details, see Fig. 4). In our model, the intron forming the 16-nt hBHBh' at canonical position 37/38 is spliced out after processing the two introns forming the 22-nt hBHBh' motif and the 26-nt HBh' motif located at 32/33 and 45/46, respectively. We propose these genes as a new type of archaeal tDNA containing three introns.

Most of the previously documented introns located at noncanonical positions of tDNAs have been found in Crenarchaeal genomes that encode heterotetrameric endonuclease subunits, and >90% of the novel tDNA candidates predicted in this study contained noncanonical introns (39 of out 42) and are encoded in three Crenarchaeal genomes (*P. aerophilum*, *T. pendens*, and *S. acidocaldarius*) with their heterotetrameric enzymes. On the other hand, the remaining three candidates (UM01, UM02, and MH01) were screened from the Euryarchaeal genomes of RC-I and *M. hungatei*, which are predicted to encode homodimeric enzymes (Table 1). Although no tDNAs containing introns at noncanonical positions have been found in genomes encoding homodimeric enzymes, and although it is still unclear whether the introns located at noncanonical sites in tDNAs are recognized by homodimeric enzymes, we suggest the existence of such tDNAs processed by homodimeric enzymes in Euryarchaea.

In summary, we developed a new program, SPLITSX, for detecting tDNAs containing one or more introns at noncanonical positions on the basis of BHB motif prediction. SPLITSX was able to identify all documented tRNA genes as well as several novel candidates. We further analyzed novel tDNA candidates complementing missing tRNAs in *T. pendens* and RC-I, and we suggested the existence of a new type of intron-containing tDNA with three introns within a single gene. Our list and the SPLITSX software will be useful for the elucidation of tRNA splicing mechanisms in the Archaea.

MATERIALS AND METHODS

Preparation of genome sequences

The genome sequences of 22 Euryarchaea, five Crenarchaea, and one Nanoarchaea were obtained from GenBank via the National Center for Biotechnology Information (NCBI) ftp server (<ftp://ftp.ncbi.nlm.nih.gov>). A draft assembly genome of *T. pendens* (belonging to the Crenarchaeota kingdom) was also used; the genome sequence was obtained from the United States Department of Energy (DOE) Joint Genome Institute (JGI) http server (<http://genome.ornl.gov/microbial/tpen/>). See Table S1 of the Supplemental Materials (http://splits.iab.keio.ac.jp/RNA_SupplMat.pdf)

for a comprehensive listing. The list of the 32 previously identified tDNAs containing introns at noncanonical positions was obtained from the literature (Marck and Grosjean 2003; Randau et al. 2005). Documented tDNAs were used as positive controls for our computational predictions.

SPLITSX

Computational approaches for searching BHB motifs were based on a sequence homology search and structural prediction. For the sequence homology search of BHB motifs, position weight matrices (PWMs) for the sequences of the two outer helices (h and h'), the central helix (H), and the bulge (B) within the 5' side of the BHB motif sequence (11 nt), and those within the 3'-side sequence (11 nt), were defined by a machine-learning approach from documented BHB motifs (Marck and Grosjean 2003; Randau et al. 2005). To calculate PWMs, the number of occurrences of each base at a given position was compiled. SPLITSX detects pairs of those 5'- and 3'-consensus sequences having lengths between 11 and 200 nt, and determines this region as the first BHB motif candidate. The first BHB motif candidates are further screened for the minimal BHB secondary structure model. The minimal structure of the BHB motif (relaxed HBh') consists of a 4-bp central helix (H) allowing a 2 bp mismatch, the 3-nt 5'-side bulge (B), and the outer helix (h') of >1 bp. The free energies of the predicted BHB structure are calculated by using RNAeval (Schuster et al. 1994) implemented within the Vienna-RNA package (<http://www.tbi.univie.ac.at/RNA/>). A cutoff score of free energy of 3 kcal/mol, which could detect all of the documented BHB motifs in tDNAs, was employed. Finally, the sequences between two bases downstream of the central helix (H) within each BHB were defined as introns.

All possible patterns of genome sequences generated with all combinations of removal of predicted tRNA introns were automatically searched using tRNAscan-SE inside SPLITSX. Because SPLITSX detects false positives within documented tDNA regions that do not contain endogenous introns, the original genome sequence without the removal of the introns was also queried to tRNAscan-SE in order to predict high-integrity candidates. If candidates from the intron-removed sequence overlapped with others, including genes annotated only by the tRNAscan-SE process, we eliminated those candidates whose COVE scores were lower than the COVE scores of their overlapping ones. tRNAscan-SE was invoked using the `-A` switch to load the specific covariance model for archaeal tDNAs. SPLITSX was performed with the following parameters: `-d 2`, `-p 0.51`, `-H 2`, and `-F 3`. The SPLITSX source code and the software package are freely available at our Web site (<http://splits.iab.keio.ac.jp>).

Comparative genomics analysis of tRNA-splicing endonuclease subunits

Using whole-genome sequences whose tRNA-splicing endonuclease subunits were still unclear and unclassified into the types of structural features, we conducted BLASTP searches, using the DNA sequences encoding the tRNA endonuclease subunits of *M. jannaschii* as the query. According to the analysis of Tocchini-Valentini et al. (2005), a species was defined to have a homotetrameric enzyme if only one homologous gene was found in the genome. Likewise, a heterotetrameric enzyme was defined to be

present when two homologous genes were observed in separate positions, and a homodimeric enzyme was defined when two homologous sequences were within the DNA region that has a translated amino acid sequence length of <350 residues and that was encoded by one gene. Finally, using MAFFT software (Kato et al. 2005), predicted amino acid sequences were aligned with known tRNA endonucleases. Both BLASTP and MAFFT were conducted by default parameters.

ACKNOWLEDGMENTS

We thank the members of MGSP at the Institute for Advanced Biosciences, Keio University, for their critical suggestions. We also thank the United States Department of Energy Joint Genome Institute (<http://www.jgi.doe.gov/>) for permission to use the sequence data of *Thermofilum pendens* Hrk 5. This research was supported in part by the Japan Society for the Promotion of Science (JSPS); a grant from the Ministry of Education, Culture, Sports, Science and Technology of Japan (The 21st Century COE Program, entitled Understanding and Control of Life's Function via Systems Biology).

Received September 21, 2006; accepted February 2, 2007.

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