

# Structural and evolutionary classification of G/U wobble basepairs in the ribosome

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## ABSTRACT

**We present a comprehensive structural, evolutionary and molecular dynamics (MD) study of the G/U wobble basepairs in the ribosome based on high-resolution crystal structures, including the recent *Escherichia coli* structure. These basepairs are classified according to their tertiary interactions, and sequence conservation at their positions is determined. G/U basepairs participating in tertiary interactions are more conserved than those lacking any interactions. Specific interactions occurring in the G/U shallow groove pocket—like packing interactions (P-interactions) and some phosphate backbone interactions (phosphate-in-pocket interactions)—lead to higher G/U conservation than others. Two salient cases of unique phylogenetic compensation are discovered. First, a P-interaction is conserved through a series of compensatory mutations involving all four participating nucleotides to preserve or restore the G/U in the optimal orientation. Second, a G/U basepair forming a P-interaction and another one forming a phosphate-in-pocket interaction are replaced by GNRA loops that maintain similar tertiary contacts. MD simulations were carried out on eight P-interactions. The specific GU/CG signature of this interaction observed in structure and sequence analysis was rationalized, and can now be used for improving sequence alignments.**

## INTRODUCTION

Comparisons of basepair frequencies and positions in conserved RNA sequences, such as rRNA, can lead to important predictive models for other RNAs. Since the sequencing of

tRNA in the 1960's, comparative sequence analysis has been used extensively to infer the common secondary structures of homologous RNAs. More recently it has been applied to infer the locations of tertiary interactions that stabilize folded RNA 3D structures. The inferred tertiary interactions in several cases have been applied to construct 3D models for biologically active RNAs, which subsequently were verified by X-ray crystallography. Recent examples include RNase P and the Group I intron (1–4). In this work we explore the potentials of tertiary interactions made by G/U wobbles to be used in similar ways.

The *cis* Watson–Crick (WC) G/U basepair is the most common non-classical basepair present in RNA. It was first recognized by Crick in 1966 in the context of the tRNA–mRNA anticodon–codon interaction (5). Subsequently, G/U's at specific positions have been shown to be essential for the function of various RNAs (6). Critical functional roles also have been inferred for highly conserved G/U basepairs found near the active sites of certain ribozymes, such as the Group I introns (4).

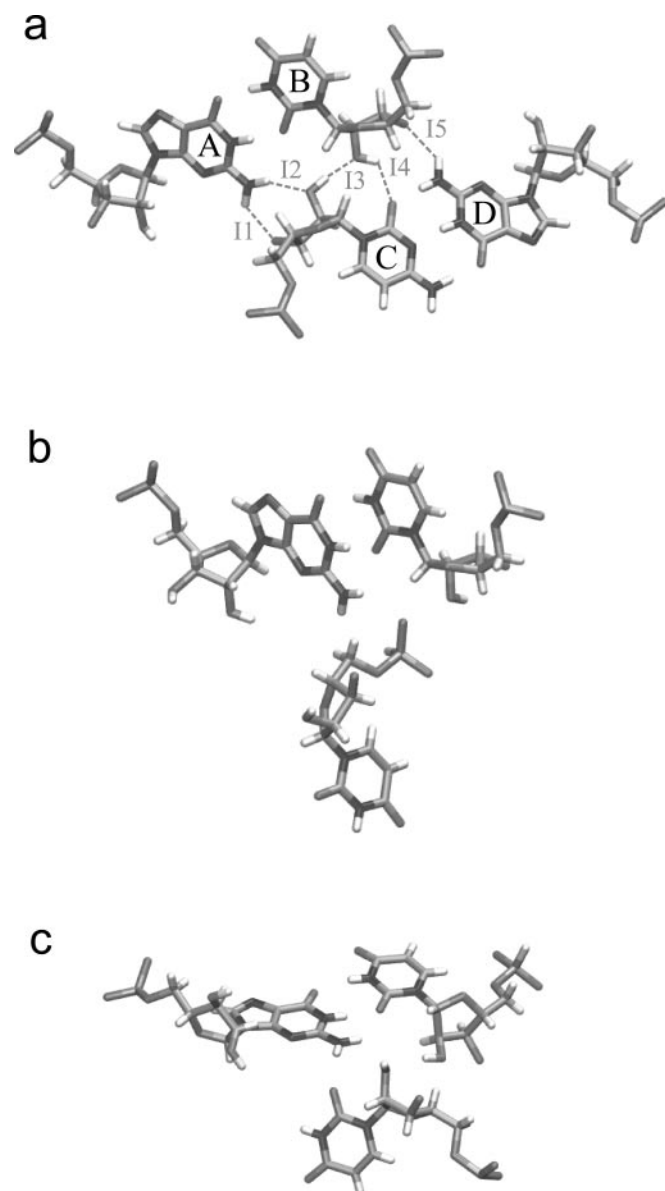
The first systematic study of the *cis* WC G/U basepairs was done by Gutell *et al.* (7) who investigated G/U variations in rRNA among broadly divergent phylogenetic taxa and classified them into several types according to their sequence conservation. In a similar way, Gautheret *et al.* (8) classified the G/U basepairs according to their phylogenetic patterns combined with their positions in the secondary structures. These authors recognized that at least 50% of WC-paired positions in 16S and 23S rRNAs contain >1% G/U in sequence alignments, and ~10% of all paired positions display 50% or more G/U substitutions. They also noted that most of these positions are often substituted by various classical WC pairs (A/U, U/A, G/C or C/G) but some highly conserved G/U's have specific variation patterns—such as G/U to U/G or G/U to A/C (8).

Earlier studies describing G/U basepairs were not based on knowledge of X-ray structures (7,8). Thus the role of tertiary interactions in constraining the G/U pairs could not be

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considered. Here, we analyze phylogenetic substitution patterns of the G/U basepairs in different structural contexts. Our work has several aims. First, to provide complete structural and phylogenetic characterization of all G/U basepairs in rRNA and classify their tertiary interactions. Second, to identify G/U-specific interactions and describe their sequence signatures. Third, to improve sequence alignments based on the tertiary interactions.

The packing interaction (P-interaction) (9) involves two basepairs, usually *cis* WC G/U and *cis* WC C/G basepairs. It somewhat resembles a 'type 0' A-minor motif (10), but with better and deeper packing of the interacting helices along the G/U shallow (minor) groove pocket (Figure 1a). Both the P-interaction and 'type 0' A-minor motif are variants of



**Figure 1.** SGP interactions made by G/U basepairs. (a) P-interaction between the 4 nt A, B, C and D. This interaction is optimal when one of the basepairs is *cis* WC G/U, resulting in the five H-bonds indicated (I1-I5). (b) Phosphate-in-pocket interaction. (a) and (b) are from PDB file 1S72, (c) is from PDB file 1J5E. Produced by DeepView (14).

ribose zippers (4). We show that P-interactions have the highest G/U conservation among all G/U interactions. We identify a novel quadruple compensatory mutation (involving 4 nt at once) between a G/U and another basepair, reinstating the interaction in a reversed orientation. We suggest that the P-interaction can be used for improving sequence alignments because of its specific GU/CG signature and, consequently, we refine the sequence alignments at several positions where a P-interaction takes place.

We identify two new types of highly conserved tertiary interactions involving the G/U shallow groove pocket (SGP). Phosphate backbone interaction (Phosphate-in-pocket interaction) has a phosphate embedded deep in the G/U SGP (Figure 1b), in analogy with the sulfate ion interaction in the same pocket (11). O2'-in-pocket interaction (Figure 1c) involves the sugar O2' coming from outside the plane of the G/U and binding at the G/U SGP.

We further show that tertiary contacts of P- and phosphate-in-pocket interactions can be conserved even upon substantial change of the local motifs involved. Thus, we found G/U's forming a P-interaction or phosphate-in-pocket interaction in some ribosomal X-ray structures to be replaced by GNRA loops in others while keeping similar tertiary contacts. These unique motif swaps underline the precedence of tertiary over secondary structure in their particular contexts.

## MATERIALS AND METHODS

X-ray crystal structures were obtained from the Nucleic Acid Database and the Protein Data Bank (PDB) (12,13) and were visualized in 3D with DeepView (14). We studied all *cis* WC G/U basepairs in the 16S rRNA of the bacterium *Thermus thermophilus* (*Tr*)—PDB files 1IBM, 3.31Å resolution, and 1J5E, 3.05Å (15), and in the 23S/5S rRNA structures of the archaeon *Haloarcula marismortui* (*Hm*)—PDB files 1JJ2 and 1S72, both at 2.4 Å (16) and the bacterium *Deinococcus radiodurans* (*Dr*)—PDB files 1KPJ and 1LNR, both at 3.10 Å (17). In addition, all *Escherichia coli* (*Ec*) *cis* WC G/U basepairs were studied in recent 70S structures—PDB files 2AVY, 2AW4, 2AW7 and 2AWB, solved at 3.5 Å (18). Structures with bound mRNA and tRNA or substrate analogues also were examined (19–26).

The alignments for the 16S-like and the 23S-like rRNA were obtained from the European ribosomal RNA database (27,28). The 16S-like rRNA alignments included 220 archaeal, 4475 bacterial and 5248 eukaryal unique sequences (sequences with only one representative from each species; thus the alignments are made of broadly divergent taxa, and their sequence analysis is less biased). The 23S-like rRNA alignments included 24 archaeal, 184 bacterial and 137 eukaryal unique sequences. The 5S-like seed RFAM rRNA alignment was used in our study (29,30). It contains 37 archaeal, 336 bacterial and 222 eukaryal unique sequences. Genomic tRNA alignments were obtained from the compilation of tRNA sequences and sequences of tRNA genes (31), and they are composed of 317 archaeal, 1768 bacterial and 141 eukaryal unique sequences. The sequence analysis was carried out with 'ribostral', a specially designed MATLAB program (A. Mokdad and N.B. Leontis, in preparation). This is a user-friendly program that includes a sequence viewer able to display substitution patterns of basepairs in an RNA alignment

colored according to their isostericity with the actual basepair observed in the crystal structure. Thus, the 3D structure information is directly related to the sequence alignment, and substitutions that belong to the same or different isosteric subfamilies for each basepair are easily recognized. Ribosomal also measures substitution patterns of multiple nucleotides at once, such as base triples or quadruples, which was useful for simultaneous analysis of all 4 nt participating in P-interactions. Further details and a preliminary stand-alone version of ribosomal can be found at <http://rna.bgsu.edu/mokdad/ribostral>.

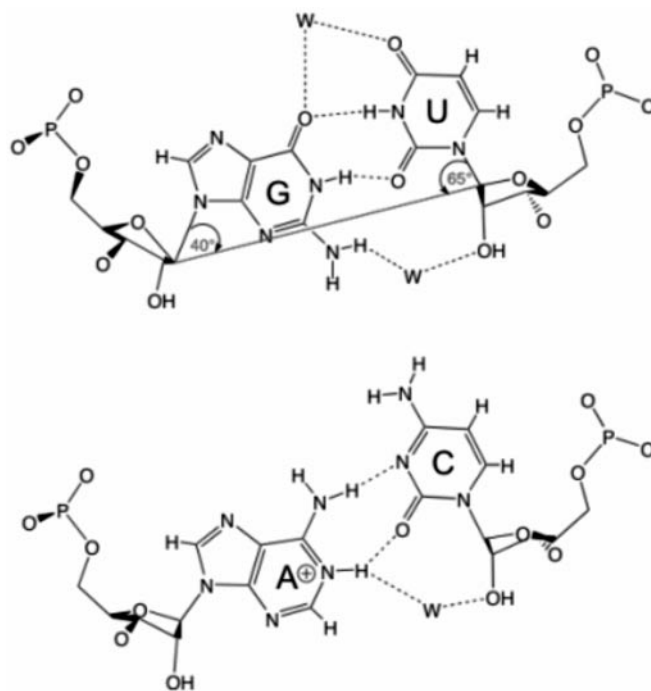
Our study includes all *cis* WC G/U basepairs seen in the high-resolution crystal structures of the ribosome and its subunits. Further, other basepairs are considered if they have >50% G/U substitutions in sequence alignments of archaeal or bacterial sequences (as these prove to be more trustworthy than eukaryal alignments). Secondary structures from the comparative RNA website (32) were first used to identify all such pairs. However, X-ray structure examination revealed that some of them were either not *cis* WC-paired or not paired at all. These basepairs were excluded from this study (for a list of basepairs omitted in this way see Supplementary Table S2). This process produced a final list of 80 bp from 16S, 186 bp from 23S and 7 bp from 5S rRNAs.

Molecular dynamics (MD) analysis was carried out on eight systems each composed of two 24 nt-long helices coming together and forming a P-interaction in their center. The first of these systems, extracted from PDB file 1S72, contains a G/U packed with C/G (P-interaction, abbreviated as GU-CG). This central P-interaction was then modified using InsightII (from Accelrys) to obtain the following eight different combinations: GU-CG (original), AC-CG, GU-GC, AC-GC, GC-CG, GC-GC, AU-CG and AU-GC. The explicit solvent MD simulations were carried out using the AMBER7.0 package (33) with the parm99 Cornell *et al.* force field (34–36). The RNA was solvated in a rectangular box of TIP3P waters (37) and neutralized by minimal number of sodium cations (38) initially placed by the Leap module at points of favorable electrostatic potential close to the RNA. The standard protocols (39) were used for the equilibration and production simulations, performed by the Sander module of AMBER7.0. The production runs were carried out at 300 K with constant-pressure periodic boundary conditions and the particle mesh Ewald method (40) applied. The MD trajectories were then analyzed using the ptraj and carnal modules of the AMBER7.0 package and our own scripts and visualized by the program VMD (41).

## RESULTS AND DISCUSSION

### Specific features of the *cis* WC G/U basepair

The C1'–C1' distance of the *cis* WC G/U basepair is  $\sim 10.2$  Å, which is very close to the distances in classical WC basepairs (10.3 Å). The U is shifted towards the deep (major) groove to allow for hydrogen bonding between G(O6) . . . U(N3) and G(N1) . . . U(O2). The angle between the C1'–C1' axis and the glycosidic bonds is  $\sim 40^\circ$  for G and  $65^\circ$  for U, instead of the symmetric  $54^\circ$  angle in the canonical basepairs (Figure 2). This causes helices containing G/U's to locally overtwist or undertwist depending on the orientation of the wobble basepair and its neighbors (6). Therefore, the only other basepair



**Figure 2.** Top, *cis* Watson–Crick G/U (wobble) basepair with water molecules (W) in its SGP and DGP. The angles formed between the C1'–C1' axis and the glycosidic bonds show the asymmetry of this basepair compared with the classical WC basepairs. Bottom, the isosteric A<sup>+</sup>/C basepair. Produced by ChemDraw (CambridgeSoft Corporation).

that is completely isosteric to the *cis* WC G/U basepair is the *cis* WC A<sup>+</sup>/C basepair, which is rare due to the A(N1) protonation requirement (42,43). The canonical WC basepairs are nearly isosteric to G/U or U/G (6) while, due to their asymmetry, G/U and A<sup>+</sup>/C are not isosteric upon reversal to U/G and C/A<sup>+</sup>. The pocket created by G(N2), U(O2), and U(O2') in the shallow groove often coordinates an integral water molecule by forming two H-bonds. Another feature of the *cis* WC G/U basepair is the unique H-bond donor and acceptor distribution around its grooves. The hydrogens of the amino group G(N2) are free to interact in the shallow groove, and the electronegative G(N7), G(O6) and U(O4) atoms in the deep groove create a strong electronegative surface (44,45) that binds metal ions. This may help in RNA folding or ribozymatic activity (46,47). In addition, the *cis* WC G/U basepair is approximately as thermodynamically stable as the classical *cis* WC A/U basepair (45,48–53). Finally, the *cis* WC G/U basepair possesses a unique conformational flexibility (6) that allows it to respond to sequence contexts and crystal packing much more easily than classical basepairs (54), hence allowing for recognition of interacting proteins or other RNAs by induced fit (55,56). These characteristics of G/U basepairs play major roles in determining the types of RNA–RNA, RNA–protein or RNA–metal ion interactions in which they participate, and in their preferred substitution patterns throughout evolution.

### Survey of G/U basepairs

Table 1 is a compilation of the secondary structure features and tertiary interactions of all the G/U basepairs that occur

**Table 1.** Interactions of the *cis* WC G/U basepairs in 16S rRNA, based on the crystal structures of *T.thermophilus* and *E.coli*

16S no.	<i>E.coli</i>			Secondary structure	Interactions	Sequence %GU+UG		
	BP	NT1	NT2			A	B	E
1	GU	22	12	h1	SGNP: A913 type1 A-minor; Mg in DGP (Tt)	7	99	1
2	GU	15	920	Part of pseudoknot	SGNP: UO3'-G1081O2' (Tt)	99	99	98
3	CG	396	45	h4	None	60	28	1
4	CG	52	359	h5, flanking IL	None	0	1	0
5/c	UG	105	62	h6	SGP: U-pack-C379 (Tt)	100	99	96
6	UA	70	98	h6	None	18	13	13
7	GU	76	93	h6	None	0	20	4
8	GC	79	90	h6	None	—	10	18
9	GU	122	239	h7, flanking 4WJ	None	0	46	1
10	AU	236	125	h7	SGNP: G-C121 tHW (Tt)	90	75	0
11	GA	232	129	h7, flanking bulge	SGNP: G-A263 tSW (Tt)	0	2	3
12	GU	226	137	h7	None	0	3	3
13	UG	157	164	h8	Mg in DGP (Tt)	100	59	54
14 (Tt)	GU	184	195	h9, flanking 4WJ	None (Position absent in Ec due to shorter helix than Tt)	4	20	0
15	GU	198	219	h10, flanking 4WJ	SGNP: G-U173 cSH (Ec)	5	13	2
16	GU	201	216	h10, flanking hairpin	None	8	73	7
17	GU	275	249	h11, flanking bulge	SGP: U252O2' (Tt)	93	99	97
18	GU	258	268	h11	None	0	7	1
19	GU	293	304	h12, flanking bulge	None	0	56	4
20/a	GU	301	296	h12, flanking hairpin	SGP: U-pack-C556 (Tt)	100	99	99
21	GU	376	387	h15, flanking IL	Mg in DGP (Tt)	27	99	3
22	GU	433	409	h16, flanking IL	None	32	42	1
23	GU	416	427	h16, flanking IL	SGNP: UO2'-G541 Phosphate, near pocket (Tt)	—	77	67
24	GU	417	426	h16	SGNP: residues 36–45 of S4 prot; GNH2-G540O2', near pocket (Ec)	—	93	2
25	GU	454	479	h17, inside IL	None	—	19	40
26	GU	474	458	h17	None in Ec, shorter helix in Tt	—	19	0
27	GC	761	580	h20, flanking IL	None	2	5	0
28/d	GU	584	757	h20, flanking IL	SGP: U-pack-C879 (Tt)	98	100	18
29	GU	650	589	h21	None	5	19	26
30	UG	593	646	h21	None	11	63	11
31	GU	645	594	h21, flanking bulge	None	47	56	10
(32)	GC	601	637	h21	None	13	75	17
33	AU	635	603	h21	None	5	7	2
34	GU	633	605	h21	SGP: G126 Phosphate (Tt)	61	91	25
35	GU	615	625	h21	None	6	31	8
36	UA	662	743	h22	None	0	26	4
37	GU	666	740	h22, flanking bulge	SGNP: Ser52 of S15 prot (Tt)	48	96	3
38	GU	734	672	h22, flanking 3WJ	None	0	53	2
39	GU	713	677	h23, flanking IL	RNA–protein bridge (B7b); SGNP: A777 type0 A-minor (Tt)	5	74	94
40	GU	683	707	h23	SGNP: Gly37 of S11 prot (Tt)	1	97	6
41	GU	778	804	h24, flanking bulge	SGNP: GO2'-Arg120O1 of S11 prot (Tt)	65	96	2
42	AU	855	831	h26	None	100	63	3
43	GU	832	854	h26	SGNP: GO2'-G724 Phosphate, near pocket (Tt)	94	87	1
44	CG	853	833	h26	SGNP: GNH2-G725 Phosphate, near pocket (Tt)	96	89	97
45	GU	852	834	h26	None	28	18	1
46	GU	851	835	h26	SGNP: GNH2-C744O2' (Tt)	1	20	1
47	GU	849	837	h26	SGNP: UO4'-G745O2' (Ec)	17	32	4
48	GU	836	850	h26	SGNP: GNH2-C745O3' (Tt)	4	86	41
49	GU	886	911	h27	DGNP: UO2P-Arg97NH2 of S12 prot; G1489O2' near SGP; Mg in DGP (Tt)	98	96	100
50	GU	894	905	h27, flanking IL	SGNP: UO2'-U244 Phosphate (Tt)	98	99	96
51	GU	895	904	h27	None	0	39	3
52	GU	925	1391	h28, flanking bulge	SGNP: GNH2-A1503 Phosphate, near pocket; Mg in DGP (Tt)	100	100	99
53	GU	927	1390	h28, flanking bulge	SGNP: GNH2-U1532 Phosphate, near pocket; Mg in DGP (Tt)	100	100	1
54	GU	942	1341	h29	SGNP: GNH2-Gln124OE1 of S9 prot (Tt)	100	100	99
55	GU	1231	950	h30	SGNP: GNH2-G971 Phosphate, near pocket; DGNP: UO4-Thr105OG1 of S13 prot (Tt)	100	80	99
56	GU	1006	1023	h33, flanking 3WJ	None	—	14	17
57	UG	1009	1020	h33	None	—	20	25
58	UA	1017	1012	h33, flanking hairpin	None	—	7	25
59	GU	1206	1052	h34	SGNP: UO2'-A1055 Phosphate, near pocket; residues 190–194 of S3 prot (Tt)	100	100	100
60	GU	1058	1199	h34, flanking bulge	SGNP: G-G1202 tSH; Mg in DGP (Tt)	99	99	99
61	GU	1074	1083	h36, flanking 3WJ	SGNP: A1101 type1 A-minor (Tt)	24	100	99
62	GU	1099	1086	h37, flanking 3WJ	None	0	95	2
63	GU	1185	1115	h38	None	0	73	38
64	GU	1184	1116	h38, flanking 3WJ	None	13	72	6

Table 1. Continued

16S no.	<i>E.coli</i>			Secondary structure	Interactions	Sequence %GU+UG		
	BP	NT1	NT2			A	B	E
65	GU	1242	1295	h41	SGNP: G-U1302 tSW (Tt)	47	90	2
66	GU	1290	1247	h41, flanking IL	None	1	63	7
67	GU	1371	1351	h43	SGNP: GO3'-Gly69CA of S9 prot; DGNP: UO4-Lys118CE of S9 prot (Tt)	66	99	70
68	GU	1486	1414	h44	RNA-RNA bridge (B3)	0	60	1
69	GU	1415	1485	h44	RNA-RNA bridge (B3)	0	96	0
70	GU	1419	1481	h44, flanking IL	RNA-RNA bridge (B5)	15	64	1
71	GU	1422	1478	h44	None	6	50	2
72	GU	1423	1477	h44	None	76	47	2
73	GU	1475	1425	h44	RNA-RNA bridge (B5 & B6)	60	38	61
74	GU	1426	1474	h44	None	30	33	4
75	GU	1438	1463	h44	None	8	31	3
76	GU	1461	1440	h44	SGNP: G-A1441 tSH (Tt)	7	85	1
77	GU	1458	1444	h44	None in <i>Ec</i> , shorter helix in <i>Tt</i>	0	13	0
78	GU	1457	1445	h44	None in <i>Ec</i> , shorter helix in <i>Tt</i>	0	67	33
79	GC	1525	1510	h45	None	0	2	0
80/e	GU	1523	1512	h45	SGP: U-pack-A768 (Tt)	91	97	98

Initials of the structure best for observing each interaction appear in parentheses in the interactions column. Basepairs with interactions are shaded. The letters (a, b, ...) in the first column after the basepair number mark the individual basepairs forming P-interactions (indicated by -pack- in the fourth column) and are also used in Table 4 and Figure 6 and referenced in the text. Basepair number (32) is not G/U in any crystal structure, but is included in the study for having >50% GU content in bacterial sequences. The last three columns indicate GU+UG (%) content at those positions in sequence alignments of archaea (A), bacteria (B) and eukarya (E). A dash (—) in these columns means that sequence alignments are all gaps (insertions) at the corresponding location. Abbreviations: h, helix; IL, internal loop; WJ, way-junction; SGP, shallow groove pocket; SGNP, shallow groove not in pocket; DGP, eep groove pocket; and DGNP, deep groove not in pocket. For the 23S/5S rRNA analysis see Supplementary Table S3.

in crystal structures of 16S rRNA (a similar compilation for tRNA and 23S/5S rRNA interactions can be found in Supplementary Table S3). Interactions are classified according to their location, i.e. in shallow groove or in deep groove. Shallow groove interactions are subdivided further into those that involve the G/U pocket (SGP), forming H-bonds with at least two groups among GNH2, UO2 and UO2', and those that do not (Shallow Groove not in Pocket, SGNP). Deep groove interactions are subdivided into those that involve the deep groove pocket (DGP), forming H-bonds with GO6 and UO4 simultaneously, and those that do not (Deep Groove not in Pocket, DGNP). In most cases, the *Tt* and *Hm* structures prove to be the best for observing 16S and 23S/5S interactions, respectively. The 23S *Dr* and 70S *Ec* structures are at slightly lower resolutions and are mainly used for comparison and for inferring motions or motif swaps, as well as observing basepairs that are not G/U's in *Tt* or *Hm*. Table 1 also displays the conservation of the G/U in sequence alignments at each of the positions. The main goal for our classification was to test whether the interactions in the G/U SGP are more specific for G/U than others occurring elsewhere around this basepair.

Among the 80 G/U basepairs that occur in the 16S rRNA, 48 (60%) are inside helices, and the remaining 32 (40%) are at the ends of helices or within other motifs. Among the 193 G/U basepairs of 23S and 5S rRNA, 122 (63%) are inside helices, and the remaining 71 (37%) are elsewhere. This agrees with the previous counts of 56% intra-helical and 44% at the ends of helices compiled by Gautheret *et al.* (8), although our criteria for finding the G/U pairs are slightly different. The *cis* WC G/U basepairs have greater probability to be found within helices compared with *cis* WC A/G basepairs, which have significantly larger C1'-C1' distance and are thus found almost exclusively at the ends of helices (57).

Of the 80 16S basepairs, 28 (35%) form SGNP interactions, 6 (8%) form interactions in SGP and 2 (3%) form DGNP

interactions. Five basepairs form intermolecular bridges with ribosomal proteins or the large ribosomal subunit (58). No interactions are observed in the DGP. Instead, this area is occupied by a cation in several cases, consistent with the high electronegativity of this site (44,45). About one-half of the 16S G/U basepairs (42 out of 80) show no evidence of any tertiary interaction in the available crystal structures (interactions with ions are not considered as tertiary interactions). Similarly, of the 193 23S/5S G/U basepairs, 62 (32%) form SGNP contacts, 21 (11%) form interactions in SGP, 4 (2%) form DGNP interactions and 3 form interactions in DGP. Seven basepairs form intermolecular bridges with tRNA (58). A cation binds simultaneously to GO6 and UO4 in several cases, and a water molecule occupies the G/U SGP in about one-third of the cases (only *Hm* crystal structure contains water molecules). Again, about one-half of the 23S/5S G/U basepairs do not form any tertiary interactions (103 out of 193).

These data demonstrate almost total agreement between the statistics of the two ribosomal subunits, showing that most tertiary interactions involving G/U occur in the shallow groove. Almost all G/U tertiary and quaternary interactions in 16S *Tt* and *Ec* and in 23S/5S *Hm*, *Dr* and *Ec* are seen at equivalent positions. Exceptions are those interactions that occur in a variable region that is either absent in one or two of the structures, or significantly different between them. Another possible reason for seeing interactions in some of the crystal structures and not all of them is the presence of a proximal kink-turn (59). When the intrinsically flexible kink-turns change between open and closed conformations, different sets of tertiary contacts become possible. Kink-turns are like hinges that allow the motion of attached helices with respect to the body of the molecule. Upon opening or closing of the kink, the motion propagates from the fulcrum of the kink to the attached helical arms (39). Since each crystal

structure shows only a static snapshot, different X-ray structures can capture the flexible K-turns in different substates. An example of this is the position corresponding to *Hm* G798/U815 in H34 (H and h symbols followed by a number denote helices in the large and small subunit, respectively). Here, a tertiary interaction is seen in the *Hm* structure but not in the *Dr* or *Ec*. Another example is the G/U basepair corresponding to *Hm* G1646/U1539 in H56. A tertiary interaction is seen in the *Hm* structure between G1646(NH2) and A1597(NH2). This interaction is possible because GNH2 can assume a pyramidal configuration due to the NH2 partial sp<sup>3</sup> hybridization (57). No equivalent interactions are seen in *Dr* or *Ec*. Note that the limited resolution of the X-ray structures may affect the appearance of some of the interactions.

### Classification of observed tertiary interactions and their sequence patterns

Three distinct types of shallow groove interactions involve the G/U pocket (SGP): P-interactions, phosphate-in-pocket interactions and ribose O2'-in-pocket interactions (Figure 1). Aside from the G/U pocket, several other types of interactions occur in the shallow groove. These include phosphate single H-bond interactions, 'type 0' A-minor motifs, 'type I' A-minor motifs (also known as *trans* sugar edge/sugar edge or tSS basepair), H-bonding to GNH2 or ribose sugars of a G/U, and other edge-to-edge interactions, the most common of which are cSS (*cis* sugar edge/sugar edge, also known as 'type II' A-minor motif) and tWS (*trans* Watson-Crick/sugar edge) interactions. There also are >20 non-specific interactions with proteins. A few G/U basepairs participate in more than one type of tertiary/quaternary interactions at once. Some others participate in interactions that look like SGNP interactions in the crystal structure, but could easily fall into the SGP category after a subtle geometrical rearrangement ('potential' SGP interactions).

Tables 2 and 3 represent sequence analysis of G/U basepairs classified into four major categories: basepairs with SGP interactions, potential SGP interactions (see above), other types of interactions and no interactions. The percentage of basepairs at each position is measured after ignoring all sequences with gaps (deletions) (i.e. they are not percentages over all sequences). For each position, however, we also report the percentage of sequences with gaps over all sequences (shaded). If no motif swaps take place between domains or classes at these locations, then this percentage of gaps can be considered a measure of the quality of alignments at the corresponding positions.

In general, we observe that when G/U basepairs are not conserved they are replaced primarily by A/C and by classical WC basepairs (Tables 2 and 3). Much less common are substitutions by U/G or C/A or other non-classical basepairs marked as 'NTs' (other nucleotides). Although A/C and C/A basepairs are relatively scarce, and thus a trend is difficult to measure, G/U's seem to get more commonly substituted by A/C's than C/A's. As mentioned above, this can be explained by isostericity (42). This type of substitution pattern is more apparent in the category of non-interacting G/U's, whereas G/U's that make SGP interactions, and in some cases those that make potential SGP interactions are much more conserved (Tables 2 and 3). This indicates that these interactions are specific for G/U basepairs. G/U mutations at such locations can disturb 3D structure which might affect the fitness of the ribosome. One exception occurs in 16S rRNA, where the G/U's with SGP interactions have 84% GU and 7% UG content in archaeal sequence alignments. Although it seems to violate the isostericity principle, it has a simple explanation. There are only 6 bp in this category in 16S, and the 7% UG observed is due to a single position, G633/U605, which is occupied by G/U in 22% and by U/G in 39% of the sequences. This basepair makes a phosphate-in-pocket interaction, which can still take place if G/U flips to U/G,

**Table 2.** Sequence analysis of *cis* WC G/U basepairs in 16S 5S rRNA

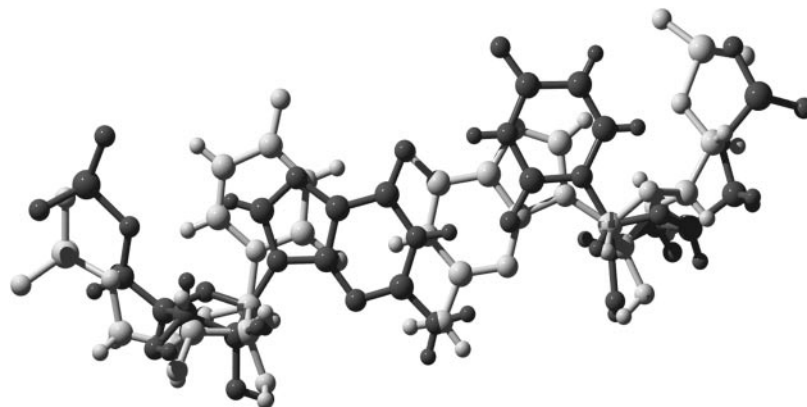
16S rRNA	Archaea (220 sequences)							Bacteria (4475 sequences)							Eukarya (5248 sequences)						
	GU	AC	WC	UG	CA	NTs	Gaps	GU	AC	WC	UG	CA	NTs	Gaps	GU	AC	WC	UG	CA	NTs	Gaps
6 interactions in SGP	84	1	4	7	0	5	10	98	0	1	0	0	2	12	68	0	14	4	3	10	34
10 potential interactions in SGP	98	0	1	0	0	0	23	92	0	7	0	0	1	2	66	11	21	0	0	2	3
22 other interactions	34	4	55	1	0	6	8	72	0	19	2	0	7	10	25	1	41	7	0	26	10
42 with no interactions	13	2	67	4	2	12	27	32	1	54	5	0	7	12	6	4	58	3	2	27	38

Percentages are measured after ignoring sequences with gaps (deletions). The percent of gaps is separately given in the last column (shaded) as a measure of the quality of alignments at the corresponding positions, assuming that no motif swaps take place. Abbreviations: SGP, shallow groove pocket; WC, any combination of the four classical Watson-Crick basepairs (G/C, C/G, A/U and U/A); and NTs, any basepair other than GU, AC, UG, CA or WC.

**Table 3.** Sequence analysis of *cis* WC G/U basepairs in 23S and 5S rRNA

23S and 5S rRNA	Archaea (24 and 37 sequences)							Bacteria (184 and 336 sequences)							Eukarya (137 and 222 sequences)						
	GU	AC	WC	UG	CA	NTs	Gaps	GU	AC	WC	UG	CA	NTs	Gaps	GU	AC	WC	UG	CA	NTs	Gaps
21 interactions in SGP	92	0	4	0	0	3	8	93	0	3	0	0	4	7	57	2	32	2	0	6	18
6 potential interactions in SGP	15	0	66	2	0	18	17	48	1	22	18	2	9	7	18	3	52	19	1	8	35
62 other interactions	37	1	55	4	0	2	3	39	1	48	2	0	7	7	23	2	56	7	1	13	11
103 with no interactions	18	2	68	3	1	7	5	30	1	53	3	1	11	7	12	3	55	5	2	22	23

Percentages are measured after ignoring sequences with gaps (deletions). The percent of gaps is separately given in the last column (shaded) as a measure of the quality of alignments at the corresponding positions, assuming that no motif swaps take place. Abbreviations: SGP, shallow groove pocket; WC, any combination of the four classical Watson-Crick basepairs (G/C, C/G, A/U and U/A); and NTs, any basepair other than GU, AC, UG, CA or WC.



**Figure 3.** G/U superimposed on U/G along the C1'–C1' axis without distorting the backbone, showing that position of the GNH2 is invariant with basepair reversal.

since position of GNH2 that makes one of the two H-bonds in this interaction remains virtually unchanged (Figure 3). This means that specific orientation of G/U is not required for the phosphate-in-pocket interaction.

The high gap count in some highly conserved regions, like those with specific SGP interactions, may suggest either mistakes in the available alignments, or motif swaps. This seems to be more apparent in the case of eukaryal alignments in both subunits. For this reason, we consider eukaryal alignments of lower quality than archaeal and bacterial alignments.

Contemporary sequence alignment methods depend almost exclusively on the local basepairing and covariation patterns of classical WC basepairs, and some G/U basepairs, and do not take into consideration tertiary interactions that impose unique constraints. Including the tertiary interaction data in sequence alignment methods would greatly enhance the quality of those alignments, and is simple to accomplish for some particular interactions with strong signatures. An example of this is 23S *Hm* U731/G740 which in crystal structures superposes with *Dr* U650/G660, both making the same P-interaction. Current sequence alignment protocols, however, do not properly align these positions. We were able in this case to use the structure data to manually adjust the alignments accordingly.

### P-interaction

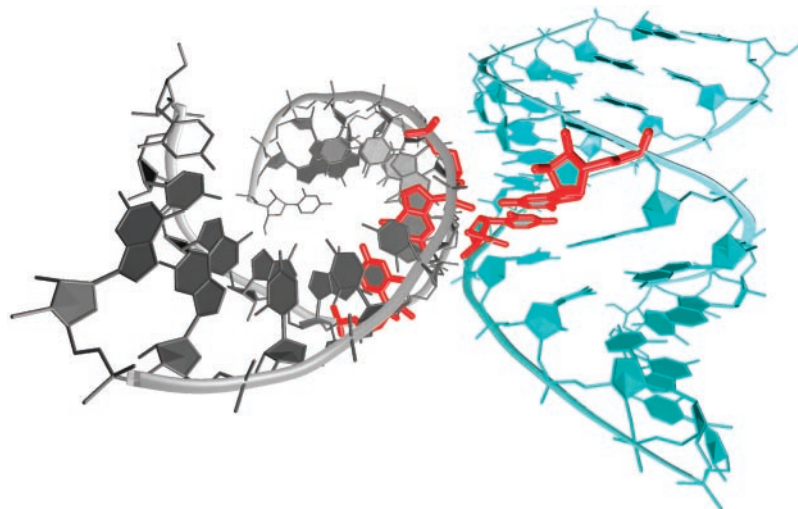
The most conserved interaction that G/U basepairs make is the P-interaction. It takes place between the ribose O2' belonging to a nucleotide from a basepair in one helix packing deep into the G/U SGP of a second helix, forming up to five H-bonds along the interface. One nucleotide from the first basepair makes most of the contact with one nucleotide from the second basepair. These 2 nt are known together as the internal pair. We designate this interaction as AB–C–D, where AB and CD are the basepairs from the two helices, with B and C forming the internal pair (as seen in Figure 1a). A previous study noted that the P-interaction plays important roles in tRNA binding to the 50S subunit and in translocation, and that it is conserved in all three phylogenetic domains of the large subunit (9). No prior sequence analysis work on the small subunit has been reported, and no attempt was made to divide the results according to the three

domains. We exhaustively searched the 3D structures for this type of interaction using several methods including FR3D, a structural motif search program of our design (M. Sarver, C.L. Zirbel, J. Stombaugh, A. Mokdad and N.B. Leontis, manuscript in preparation). Five such interactions are found in the small ribosomal subunit and thirteen in the large subunit, as reported in Tables 4 and 5. The tables include locations and substitution patterns at all four positions participating in this interaction. Among the thirteen P-interactions of the large subunit there is one between 23S and 5S, one between 23S and E-site tRNA, and one between 23S and P-site tRNA. U–C is the most commonly seen internal pair, but there is no apparent preference for a Y–Y over Y–R interactions, since U–G, U–A, and other Y–R also are seen. There is, however, clear preference for these over R–R interactions (none of these are observed). G/U on the other hand is the most commonly seen basepair making this interaction, as only 3 of the 18 P-interactions do not involve a G/U (P-motifs denoted as b, I and J in Tables 4 and 5). In addition, two such interactions involve a non-WC basepair (e and 5S). A total of four P-interactions were not reported before (e, B, I and 5S).

The high conservation of G/U's participating in P-interactions is remarkable (Tables 4 and 5). Even in the cases where the crystal structure did not contain a G/U basepair (b, I and J), alignments of some domains displayed a very high G/U content in the first helix or U/G in the second helix (95% U/G in the second helix in archaeal alignments for b, 100% G/U in the first helix in bacterial sequences for I, and 88 and 98% G/U in the first helix in archaeal and bacterial sequences for J). Even more striking is that in the rare cases when the G/U in a P-interaction is lost, there is great tendency for all 4 nt forming the interaction to mutate in a way to finally recreate a G/U in the right orientation in either one of the two involved helices. This is best demonstrated by the P-interaction corresponding to *Hm* G684/U662–C748/G657 (C in Table 5, between H28 and H27). It seems that a quadruple compensatory mutation between archaea and bacteria has occurred. The GU–CG in *Hm* is replaced by a GC–UG in *Dr*, and by CG–UG in *Ec*. In archaeal sequence alignments, these positions are 21% GU–WC, 71% WC–UG and 8% WC–WC. The latter number could correspond to organisms that survived with the intermediary mutation. Almost all (99%) of bacterial sequences are







**Figure 4.** P-interaction between two helices showing the whole system used for MD analysis; the nucleotides forming the P-interaction are highlighted in red. Produced by VMD (41).

WC–UG, and most eukaryal sequences (85%) are GU–WC. The most parsimonious explanation is that originally there was a WC–UG variant, which is still present in bacteria today. Sometime after the archaeal/eukaryal line diverged from bacteria a mutation occurred, causing some archaeal organisms to have WC–WC at those positions. Some of these organisms live till today. However, most archaea had other compensatory mutations following the first one causing either their return to the original WC–UG bacterial variant, or their ‘flipping’ into the opposite GU–WC variant. Some archaea also may have retained the original bacterial variant without ever mutating because they may have originated before the first mutation ever occurred. Eukarya most probably separated from archaea before the first mutation. These observed patterns of substitution strongly suggest that the P-interaction was conserved in most, if not all organisms, even when the identity of the individual nucleotides making it had changed. It also is clear that the G/U basepair with U at the internal position was favored. This is the first time that such a covariation involving 2 bp, rather than just 2 nt, has been reported.

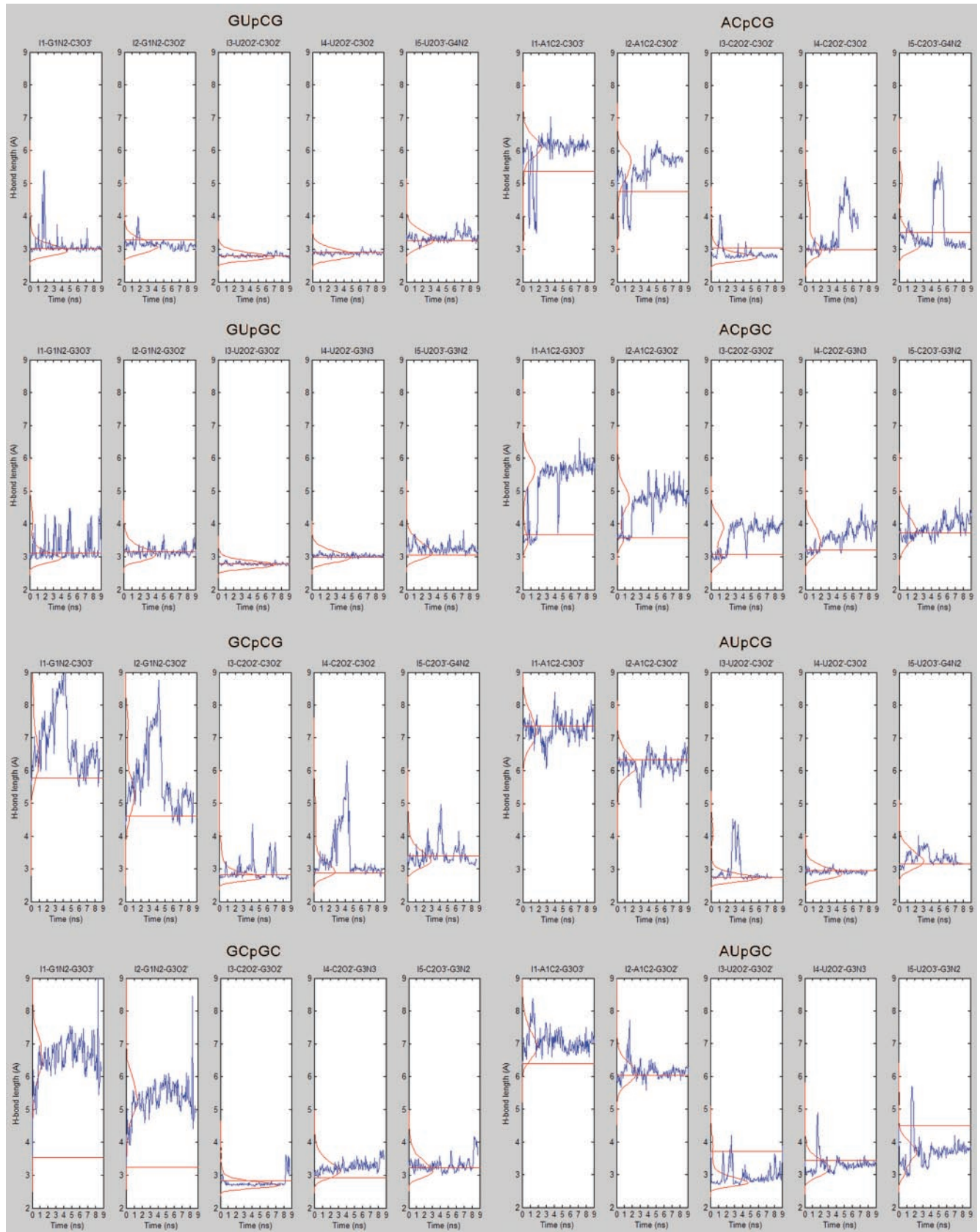
There is, on the other hand, one case where a P-interaction in bacteria (corresponding to *Ec* numbers G1878/U1864–C414/G2409) is completely lost and replaced by a GNRA loop interaction in archaea (and probably in eukarya as well, as suggested by their sequence alignments). The GNRA loop substitutes for one of the two helices (H68 flanked by the G/U) without much distortion of structure, because it makes a similar 3D contact as the P-interaction (Supplementary Figure S1). This proves that the presence and orientation of an interaction (which defines the 3D folding) is far more important than the actual identity or type of that interaction.

Tables 4 and 5 also shows that no G/U was substituted by U/G (in the same helix), which can be explained first by the fact that *cis* WC U/G is not isosteric to G/U, and second by the fact that the P-interaction, unlike the phosphate-in-pocket interaction, is structurally directional. It cannot occur as UG–CG because of the specific asymmetric orientation of the G/U basepair that is more open for SGP interactions coming from the side of the U rather than the side of the G.

### Molecular dynamics analysis of the P-interaction

Most of the P-interactions observed in crystal structures are between a G/U pair and a C/G pair, with U and C being at the internal positions (GU–CG). The purpose of MD analysis was to explain this behavior, as well as reveal why the G/U’s forming P-interactions almost never are substituted by the isosteric A/C basepairs throughout evolution.

MD simulations were carried out for eight different combinations of P-interactions, namely GU–CG, AC–CG, GU–GC, AC–GC, GC–CG, GC–GC, AU–CG and AU–GC. The studied motifs were embedded in two A-type helices, each with 24 residues (Figure 4). The helices were identical in all simulations, except for their centrally positioned P-interactions. The above combinations were chosen to represent Y–Y and R–Y types of interaction, as well as G/U and other isosteric (A/C) or nearly isosteric (A/U and G/C) basepairs embedded in one helix, combined with G/C and C/G basepairs embedded in the other helix. The simulations were extended to ~10 ns each, which appears to be sufficient for the purpose of this study. Figure 5 shows time development of the five H-bonds constituting the P-interaction in each system (as defined in Figure 1a, except that H-bond lengths are measured via the heavy atoms in the simulations). The eight structures were ranked in order of decreasing stabilities which was indirectly estimated based on the number of H-bonds and their dynamic behavior in the simulations. (Note that such classification is more relevant to judge the stability of RNA interactions than, for example, evaluation of direct base–base interaction energies. This would neglect the interplay between that base–base interaction and all the other effectors such as adjacent basepairs, solvent screening and the like). The suggested stability order is as follows: GU–CG = GU–GC (these complexes maintained all five H-bonds throughout the simulations) >> GC–GC = AU–GC = AU–CG = GC–CG (these lost the first two H-bonds, but generally maintained the others) > AC CG = AC GC (most H-bonds lost early in the simulations). Thus, a P-interaction involving A/C is the weakest among the tested systems. This is related to the fact that the A/C SGP is



**Figure 5.** MD of the eight P-interaction systems studied. All five H-bonds involved in the generic P-interaction are monitored (as defined in Figure 1a, but here are measured from the heavy atoms). The red horizontal lines mark H-bond lengths in starting structures (average over first 50 ps); the red curves describe the probability distribution of H-bond lengths in the simulation and blue curves displays the actual time development of the H-bond lengths along the trajectory.

less electronegative than the G/U pocket, and it has a much lower H-bonding potential due to the loss of NH<sub>2</sub> group. Hence, it cannot accommodate the ribose O2' in the same way as G/U does (for a comparison of electrostatic potential between G/U and A<sup>+</sup>/C see Supplementary Figure S2). Supplementary Data further present free energy calculations (MM-GBSA and MM-PBSA methods, Supplementary Table S1) of relative stabilities of the studied P-interactions. The calculations were also performed for UG-ϕ-GC combination, which can be considered as unsuitable for P-interaction. These calculations confirm superior stability of the GU-ϕ-CG = GU-ϕ-GC motifs and provide some additional insights. One has to point out, however, that such thermodynamics calculations are presently based on substantial approximations and that is why we prefer to assess the simulations primarily based on the structural dynamics seen in Figure 5.

Thus, since GU-ϕ-CG is the most commonly seen P-interaction in crystal structures and sequence alignments, and also among the two most stable systems in our simulations, it can be considered as the signature of this interaction.

### Other shallow groove pocket interactions

There are two additional SGP interactions in the ribosome, phosphate-in-pocket interaction, and O2'-in-pocket interaction (Figure 1). The latter means that O2' is inserted in the SGP of G/U, but it can not be classified as P-interaction or 'type 0' A-minor interaction. Both of these SGP interactions lead to high conservation of the G/U's. The phosphate-in-pocket interaction is less directional than the P-interaction, having no preference for a specific G/U orientation. This G/U to U/G covariation is also the case of some other interactions in the shallow groove, especially some single H-bond interactions. Thus, it is not unique to phosphate-in-pocket interactions but at least it is one of its indicators. Similar to P-interaction, we detected one case where a G/U forming a phosphate-in-pocket interaction is replaced by a GNRA loop, with no substantial distortion of the 3D folding. This occurs at the positions corresponding to *Ec* G1740/U1720-C1550P, bringing together H63 and H56. Here, both *Hm* and *Dr* have a shorter H63 than *Ec*, so the GNRA loop that normally is at the end of H63 is brought to a position homologous to the G/U position in *Ec* (Supplementary Figure S1).

Figure 6 and Supplementary Figure S3 summarize positions of all SGP interactions on the secondary structures of the small and large ribosomal subunits, respectively.

There are 273 positions in the ribosome occupied by *cis* WC G/U basepairs in one or more of the available crystal structures. This corresponds to roughly 15% of all ribosomal basepairs. About half of these are involved in tertiary or quaternary interactions. Therefore, they have important roles in the assembly and 3D folding of the ribosome and its subunits. Figure 7 is a Venn diagram representing the sequence data from archaeal and bacterial alignments mapped to the structural data of each of the positions. This provides valuable insights into the possible functions and mechanisms for some specific families of interactions. The diagram clearly shows that G/U's forming specific SGP interactions are most highly conserved. These are the P-interactions, phosphate-in-pocket interactions, and ribose O2'-in-pocket interactions. Other tertiary interactions also are included in

the figure, showing less G/U conservation in most cases. Protein interactions were discussed in detail in (60) and are not fully addressed here.

### Conservation patterns of potential SGP interactions

As noted above, potential SGP interactions are those that could form SGP interactions after a modest structural rearrangement (Tables 2 and 3). The substitution patterns in Tables 2 and 3 suggest that most potential SGP interactions in 16S actually form SGP interactions. They have high G/U conservation and phylogenetically they behave as actual SGP interactions. In contrast, most such 23S basepairs are not expected to form SGP interactions because of their low G/U conservation, except for *Ec* 2514-2570.

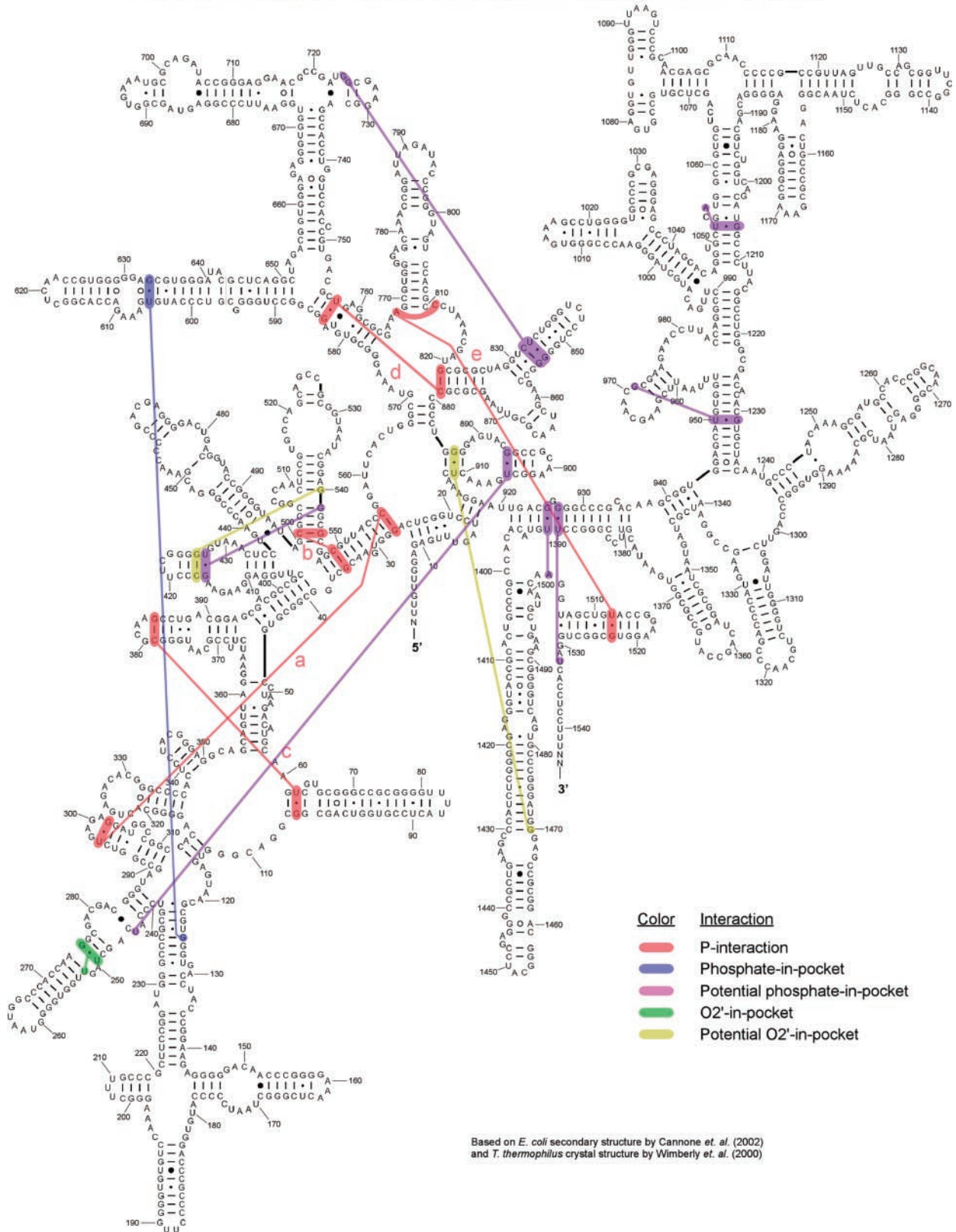
### G/U interactions can mediate flexible and transient contacts in the ribosome

The ribosome is a large dynamical molecular machine. Movements of its parts must be well coordinated for correct and efficient functioning. Transient interactions, like those with tRNA or between the ribosomal subunits, have to be formed and broken easily, and therefore must be relatively weak while still stereochemically precise. This may be possible through the association and dissociation of multiple weak tertiary and quaternary interactions or via consecutive transformation between different types of similar interactions. The similarity between P-interactions, 'type 0' A-minor motifs, O2'-in-pocket interactions, and some potential SGP interactions (all having an O2' buried at different depths in the G/U SGP) enables smooth switches from one interaction to another. A similar scenario was suggested for tertiary interactions comprising two or more A-minor motifs (61). Such dynamic behavior could explain why some G/U basepairs without apparent interactions are conserved. A possible example of this kind is 16S G886/U911 interacting with G1489/C1411, which is observed as a potential O2'-in-pocket interaction. However, sequence analysis shows that this position is >98% GU-WC (mainly GU-GC and GU-CG) in all domains, making it very similar in its evolutionary conservation to a P-interaction. We suggest that as a P-interaction relaxes or starts to dissociate, it may convert to an O2'-in-pocket interaction or a 'type 0' A-minor interaction because of the orientational similarity between them.

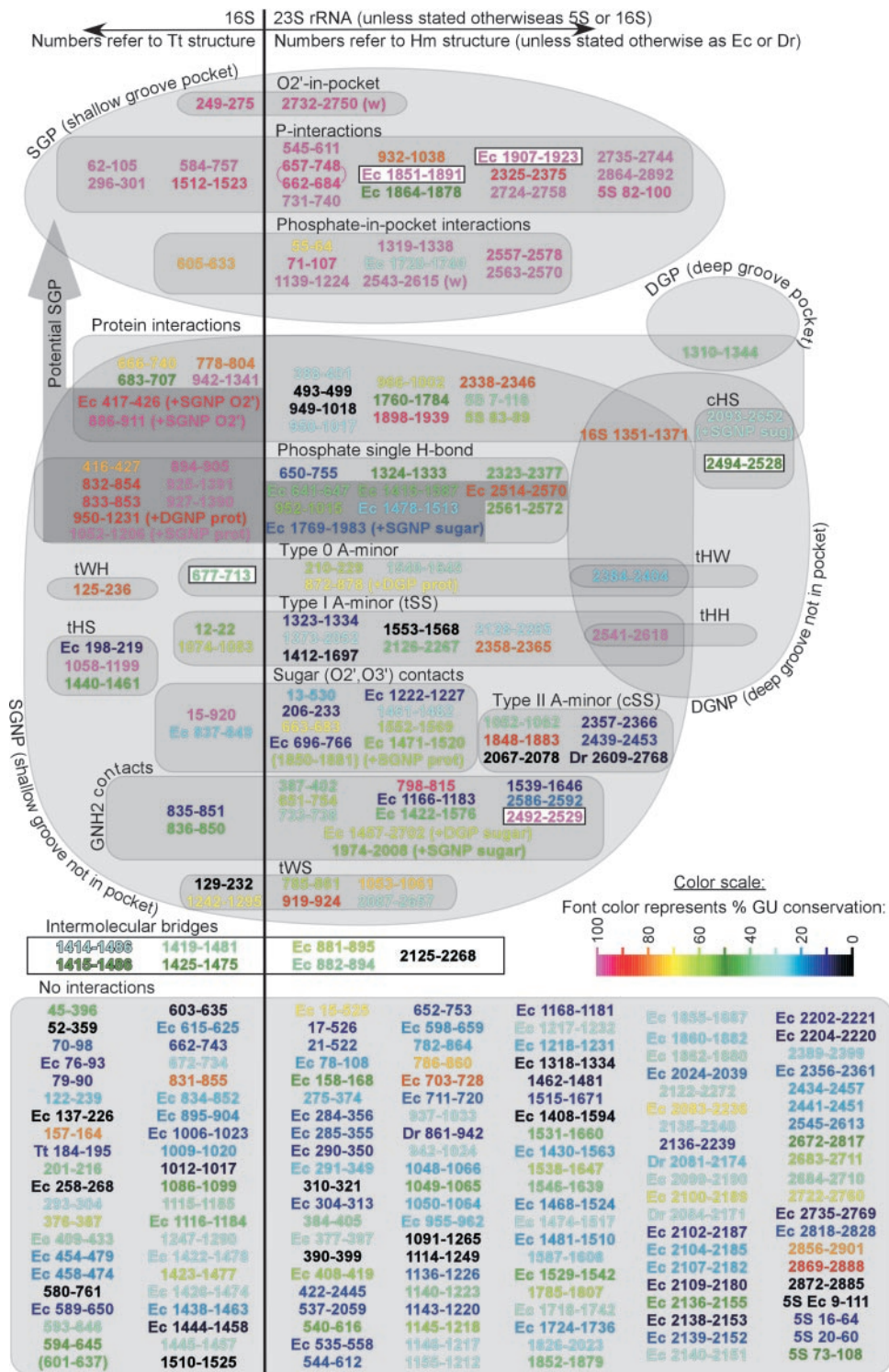
### Other conserved G/U basepairs

A few other basepairs are highly conserved in archaeal and bacterial alignments, but are not taking part in SGP or potential SGP interactions, or are not interacting at all (Figure 7). Some of these may be involved in transient interactions that are in their dissociated states in crystal structures. Others, especially those flanking internal loops or junctions which make up more than one-third of all G/U's, may be highly conserved because of the important roles they play in providing specific stacking interactions that stabilize these nearby motifs. Examples of these are 23S 1848-1883 flanking a four-way-junction at the base of H66, 23S 1898-1939 flanking the internal loop of H68, which is a conserved motif similar to C-loop, and 23S 2869-2888 flanking the internal loop of H101. Other intriguing cases include 23S 2541-2618, which is a specific basepair forming 'type I' A-minor motif with the terminal A76

Shallow groove pocket interactions (or potential interactions) in *Thermus thermophilus* 16S rRNA (numbers are based on *Escherichia coli*)



**Figure 6.** SGP interactions marked on the 16S rRNA secondary structure. Labels of P-interactions correspond to Tables 1 and 4. Color code displayed at the bottom right. For the tRNA and 23S/5S rRNA analysis see Figure S3.



**Figure 7.** Venn diagram representing locations and types of interactions of ribosomal *cis* WC G/U basepairs and their sequence conservations. Nucleotide numbers are taken from the main crystal structure in each category (*Tt* in 16S, *Hm* in 23S and 5S), unless otherwise specified (by *Dr* or *Ec* preceding the numbers). Each basepair is colored according to its average % GU+UG content in sequence alignments of archaea and bacteria. Water mediated interactions are indicated by (w); open boxes indicate intermolecular bridges between ribosomal subunits. The following abbreviations are used: c, *cis*; t, *trans*; W, Watson-Crick; H, Hoogsteen; S, sugar edge (64); SGP, shallow groove pocket; SGNP, shallow groove not in pocket; DGP, deep groove pocket and DGNP, deep groove not in pocket; additional interactions, if present, are noticed in parentheses after plus (e.g. '+SGNP sugar' and '+DGP prot' mean additional SGNP involving G/U sugar and DGP involving protein, respectively). The two positions connected by curved lines in P-interactions (657-748 and 662-684) represent the P-interaction with quadruple compensatory mutation, discussed in detail in the text. The two basepairs in parentheses (16S 601-637 and 23S 1850-1881) are not GU in any crystal structure, but are included in the study for having >50% GU content in bacterial and archaeal sequences, respectively.

of the tRNA CCA acceptor stem. The G/U here permits the interaction to become more compact by allowing the unstacked A76 to insert deeper in its pocket, making the G/U the preferred basepair at this location. The 23S 798–815 bp flanking the internal loop of H34 is another interesting case where GNH2 forms an H-bond with A1598O2', which belongs to the hairpin of H58. This helix is joined to the rest of 23S subunit by a kink-turn, and the G/U here may be involved in stabilizing conformational changes at this flexible region located at the interface with the small subunit. Another conserved G/U is 23S 2492–2529 which is a part of a complex motif forming a receptor for a GNRA loop.

## CONCLUSIONS

Large functional RNA molecules, like the ribosomal RNAs, are compactly folded and form complex tertiary and quaternary RNA–RNA and RNA–protein contacts mediated by several types of recurrent motifs and interactions. We have carried out a complete structural and sequence analysis of the G/U basepairs in rRNA and classified their tertiary interactions.

The SGP interactions of G/U basepairs reaching deep in shallow grooves of helices are clearly the most prominent G/U interaction patterns. They include the P-interaction identified earlier (9) and phosphate-in-pocket interactions and O2'-in-pocket interactions reported here for the first time (Figure 1). We also detected several P-interactions not noticed before. We show that the P-interaction G/U's are the most conserved ones, closely followed by the remaining SGP interactions.

We identify a novel quadruple compensatory mutation (involving 4 nt at once) between a G/U and another basepair, reinstating the P-interaction in a reversed orientation. Further, we show that tertiary contacts of P- and phosphate-in-pocket interactions can be conserved upon considerable change of the local motifs involved. Thus, one G/U forming a P-interaction in *E.coli* and *D.radiodurans* is replaced by a GNRA loop in *H.marismortui*. Still, similar tertiary contacts are present between the equivalent areas of the structures. Another G/U making a phosphate-in-pocket interaction in *E.coli* is replaced by a GNRA loop in both *H.marismortui* and *D.radiodurans* while again keeping similar tertiary contacts, preventing any major differences in the overall folds. These unique motif swaps underline the precedence of tertiary over secondary structure in their particular contexts. All these findings also clearly demonstrate the importance of G/U SGP interactions in folding and function of ribosomes.

We also identify interactions that could fall into the SGP category after a subtle geometrical rearrangement and we call them potential SGP interactions. This could indicate a transient formation of SGP interactions in these positions. Indeed, most potential SGP interactions in 16S rRNA actually appear to form SGP interactions because phylogenetically they behave as SGP interactions. In contrast, most such 23S rRNA basepairs are not expected to form SGP interactions because of their low G/U conservation. The similarity between P-interactions, 'type 0' A-minor motifs, and O2'-in-pocket interactions, and some potential SGP interactions indicates that they could be involved in smooth structural switches utilizing a set of consecutive similar interaction patterns.

Most of these G/U interactions are long-range contacts bringing together different RNA molecules or distant helices in the same RNA molecule. Some of them, however, are short range interactions that are nonetheless extremely conserved, such as the P-interactions b, C, D and J, the phosphate-in-pocket interactions occurring in h34, between H6 and H7, and in H91, and the O2'-in-pocket interaction occurring in h11 (refer to Figure 6 and Supplementary Figure S3). Interestingly, none of these interactions takes place between the 'head' of the 30S subunit corresponding to domain III in 16S rRNA and the rest of the small subunit. This agrees with previous studies showing that only one A-minor motif interaction occurs between domain III and the rest of the small subunit (61). It also supports the observation in the 70S *Ec* that the 'head' is subject to large-scale rigid body motions relative to the rest of the ribosome during the protein synthesis cycle, especially the ratchet-like motion that accompanies translocation (18,62,63).

In contrast to SGP interactions, G/U DGP interactions are very rare. There are just three occurrences of these in both ribosomal subunits. These occurrences do not resemble each other structurally, preventing their classification and characterization. They also do not show any specific sequence signatures, and the G/U's forming them are not highly conserved in general (Figure 7).

As noted above, few G/U's participating in P-interactions and phosphate-in-pocket interactions, which are some of the most conserved interactions in RNA, can be replaced by other elements (like GNRA loops) that are able to form similar tertiary contacts. This proves that these tertiary contacts are essential for the survival of the organisms. This, however, does not challenge the importance of the G/U basepair itself since it still remains the preferred moderator of such interactions. This knowledge can be helpful in refining sequence alignments by looking for the signatures of some of these interactions, such as the GU- $\diamond$ -CG signature of the P-interaction, and the G/U covariation with U/G which is one of the indications of a phosphate-in-pocket interaction.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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*Conflict of interest statement.* None declared.

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