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Distribution of Hammerhead and Hammerhead-like RNA Motifs Through the GenBank

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Hammerhead ribozymes previously were found in satellite RNAs from plant viroids and in repetitive DNA from certain species of newts and schistosomes. To determine if this catalytic RNA motif has a wider distribution, we decided to scrutinize the GenBank database for RNAs that contain hammerhead or hammerhead-like motifs. The search shows a widespread distribution of this kind of RNA motif in different sequences suggesting that they might have a more general role in RNA biology. The frequency of the hammerhead motif is half of that expected from a random distribution, but this fact comes from the low CpG representation in vertebrate sequences and the bias of the GenBank for those sequences. Intriguing motifs include those found in several families of repetitive sequences, in the satellite RNA from the carrot red leaf luteovirus, in plant viruses like the spinach latent virus and the elm mottle virus, in animal viruses like the hepatitis E virus and the caprine encephalitis virus, and in mRNAs such as those coding for cytochrome P450 oxidoreductase in the rat and the hamster.

The hammerhead ribozyme originally was discovered as a self-cleaving motif in viroids and satellite RNAs. These RNAs replicate using the rolling circle mechanism, which generates long multimeric replication intermediates. They use the cleavage reaction to resolve the multimeric intermediates into monomeric forms. The region able to self-cleave has three base paired helices (I–III) connected by two conserved single stranded regions and a bulged nucleotide (Forster and Symons 1987; for reviews see Symons 1992; Bratty et al. 1993; Birikh et al. 1997). The hammerhead ribozyme also seems to function in the generation of unit length sequences from multimeric transcripts of repetitive DNA sequences. Two of these RNAs have been characterized: one in several newt species (Epstein and Gall 1987) and the other one in three Schistosoma species (Ferbeyre et al. 1998). Among the repetitive sequences of these two organisms, note that not all contained a bona fide hammerhead ribozyme. Indeed, many mutations also were found creating variants of the original motif. Overall, the rather limited distribution of this motif contrast with the simplicity of its secondary structure in which only a core of 14 nucleotides is absolutely required for cleavage.

We recently have conducted an extensive research of different RNA motifs in the GeneBank database (Bourdeau et al. 1999). The results showed that most of

the motifs were distributed randomly among gene sequences suggesting that most RNA motifs originate by random drift. We now wish to extend these observations to the self-cleaving hammerhead ribozyme and its variants in which either an essential nucleotide in the single strand positions is allowed to be random or the identity of a conserved base pair from helices II and III is changed. We found that most of the hammerhead motifs are apparently underrepresented among gene sequences, but this comes from the bias of the GenBank for sequences with low CpG representation. We also report the finding of intriguing motifs in several repetitive sequences and mRNAs.

RESULTS

Searching for Self-cleaving RNA Motifs of the Hammerhead Type in the GenBank

The hammerhead ribozyme can be described by three helices separated by three single stranded regions of conserved nucleotides. There are three equivalent conformations of the self-cleaving hammerhead depending on which helix bears the 5' and 3' end of the motif. We named them HH-I, HH-II, and HH-III (Figure 1). The descriptors composed as input for the search program are presented beside each motif and described in the legend of Figure 1 (see also Methods). They were designed to detect any sequence with all the minimal nucleotide requirements to have some catalytic activity and with the possibility to fold like the hammerhead. In this context, it is expected that sequences will be found that combine several nonoptimum features

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identity of each one of the 2 conserved base pairs of the hammerhead motif (also boldface in Fig. 1).

Table 2 presents the data on the distribution of the mutated variants of HH-I, HH-II, and HH-III from the single stranded region. It is expected that every mutant will increase the frequency of occurrences by a factor of four because we changed the requirements in every position from only one to all four nucleotides except in position 4 where C and U already were allowed and in the cleavage site where only G originally was excluded. Thus, in position 4 we expected to double the frequency, and in the cleavage site we expected a 25% increase. The results are mostly those anticipated based on these calculations. However the mutants of position 12 doubled the expected increase in all the orientations. This effect was not uniformly observed in the different subdivisions of the GenBank. Actually, most of the extra occurrences are located in the files containing ESTs (Expressed Sequence Tags) and mammalian sequences. These preferences were not observed in the random database in which the mutants showed the anticipated increase in their frequency in comparison with the original motif. The number of occurrences obtained in the virus section of the GenBank for the HH-III-8 variant was 722 instead of the 113 expected (HH-III has 3774 expected occurrences and viruses represent 3% of the GenBank). However, a quick analysis of the occurrences obtained with this descriptor revealed that most of them are the same motif repeated in 679 hepatitis C sequences.

Table 3 presents the frequencies obtained with the mutant hammerhead ribozymes using a different identity for the conserved base pair of helices II or III (positions 10.1:11.1 and 15.1:16.1, respectively). One striking observation is that all the mutants in Helix II (iiNN) have total occurrences two to six times higher than expected whereas the mutants in Helix III (iiiNN) have half the expected frequency. One more interesting point is the high number of occurrences obtained with the three orientations of the hammerhead ribozyme having a A:U base pair in Helix II (10.1:11.1) instead of the usual G:C.

The mutants in position 12 and the mutants of the conserved base pair of Helix II have in common that they disrupt the presence of a dinucleotide CpG in the resulting sequence. It is well known that CpG is underrepresented in vertebrate sequences (Karlin and Mrazek 1997). The GenBank is biased for those sequences mainly owing to human and rodent entries. In those files, the mutants that disrupt the CpG requirement have a higher frequency. To confirm that the overall frequency of the hammerhead motifs containing CpG dinucleotides is half of the expected one because of the low CpG content of vertebrate sequences, we built a new random database in which the frequency of CpG was reduced by half in favor of either

AG, CA, CC, CT, GG, or TG to simulate the frequencies observed by Karlin and Mrazek (1997; see Methods). In this database, we observed an overall doubling of the original expected frequencies for all the motifs needing a CpG but not for the others (data not shown).

Still, the mutants with a A:U base pair in position 10.1:11.1 of the Helix II have a very high frequency in all three conformations of the motif: two to three times higher than expected even considering the CpG effect discussed above. So far, we have no explanation for this intriguing observation.

Finally, we made three more searches by changing the cleavage site from NUH to NHH based on the report of Kore et al. (1998) that such hammerheads were still active. We obtained for these new mutants a number of occurrences corresponding to half of what we expected according to the search in an equal A-C-G-T random database. Moreover, as for the previous motifs, the number of occurrences in the GenBank is comparable to the expected frequency according to the search in the reduced for CpG database. All the occurrences found in the GenBank are available in our web site at <http://www.centrcn.umontreal.ca/~bourdeav/HH>.

Some Intriguing Hammerhead Motifs that Might Have Functional Significance

This section presents a sample of motifs considered interesting either because of their location or because their structure is optimal for self cleavage. The hammerhead ribozyme occurs naturally in satellite RNAs, viroids, and transcripts from repetitive sequences. The probability of finding an active hammerhead should be higher among these genetic elements. Several potential hammerhead motifs were found in distinct families of repetitive DNA.

Hammerhead ribozymes were found in the satellite DNA from *Dolichopoda schiavazzii* (cricket) by using the HH-I descriptor (example in Figure 2A). Fourteen have a conserved HH-I motif and two have a HH-I-iiGU motif (G:U in position 10.1:11.1 instead of G:C). This ribozyme cleaves after CUA (A.A. Rojas, A. Vazques-Tello, G. Ferbeyre, F. Venanzetti, L. Bachmann, B. Paquin, and R. Cedergren, in prep.). Helix I has the GG:CC base pairs and the internal loop common to the hammerhead motifs in schistosomes (Ferbeyre et al. 1998) and newts (Pabon-Peña et al. 1991). It is noteworthy that among the 20 similar sequences submitted to GenBank, the four sequences not found through the search contained either mismatches in one of the helices or combined two point mutations.

A hammerhead-like motif was detected in the Kpn-13 family of human repetitive DNA by using the descriptor HH-I-4 (Fig. 2B). The motif is found in several ESTs containing Kpn-repetitive sequences (also known as L1-repetitive elements) indicating its expression at the RNA level. All the occurrences contain a

Table 2. Distribution of Hammerhead and Hammerhead-like Motifs in the Different Sections of the GenBank: Mutants of the Single Stranded Regions

	pri	rod	mam	vrt	inv	pln	bct	rna	vrl	phg	syn	una	est	pat	sts	gss	htg	Total	Total	Expected
																		over	expected	(A = C = G = T)
HH-I	51	21	0	14	176	119	130	0	37	3	5	0	186	45	4	17	108	916	0.56	1635
HH-I-3	182	72	12	45	997	560	553	2	159	16	6	0	702	125	10	119	486	4046	0.57	7081
HH-I-4	270	29	0	32	365	228	226	10	69	3	8	1	358	55	4	33	272	1963	0.55	3576
HH-I-5	310	70	14	97	506	400	399	2	77	10	7	1	641	60	13	92	382	2991	0.46	6488
HH-I-6	197	60	17	37	516	312	309	3	107	5	8	1	700	93	11	67	288	2821	0.49	6973
HH-I-8	171	44	11	22	790	407	455	10	138	11	5	0	521	86	10	61	420	3162	0.46	6883
HH-I-9	247	56	12	29	508	260	454	0	107	9	7	3	536	99	9	65	344	2745	0.38	7314
HH-I-12	946	224	41	103	754	613	454	0	180	13	8	2	1856	137	48	263	794	6436	0.94	6865
HH-I-13	209	76	16	41	428	308	462	5	93	10	11	0	739	143	12	66	226	2845	0.42	6775
HH-I-14	317	94	30	36	404	377	487	7	104	5	30	5	737	102	8	77	327	3147	0.50	6362
HH-I-17	65	25	0	18	201	144	162	0	50	4	4	0	225	53	4	19	122	1103	0.54	2031
HH-II	83	33	1	5	175	126	141	1	80	1	1	3	435	30	4	8	91	1218	0.54	2246
HH-II-3	180	78	8	29	994	645	715	3	204	14	25	23	1022	102	11	82	502	4637	0.66	7063
HH-II-4	133	56	7	11	387	261	278	1	100	1	2	5	658	43	7	39	188	2177	0.51	4277
HH-II-5	244	82	16	37	573	477	372	7	141	10	4	4	910	70	12	71	362	3392	0.39	8788
HH-II-6	234	81	23	17	522	348	977	38	153	7	7	9	856	74	8	59	311	3724	0.49	7674
HH-II-8	209	64	11	22	921	385	519	16	130	3	10	4	1265	53	7	65	452	4136	0.54	7710
HH-II-9	255	58	8	24	457	319	540	6	140	9	11	5	884	82	10	51	281	3140	0.39	7979
HH-II-12	1290	253	61	60	879	716	534	1	209	14	18	3	2491	125	56	273	1027	8010	0.97	8285
HH-II-13	214	91	45	26	421	326	534	7	162	8	23	3	1031	71	7	53	228	3250	0.44	7404
HH-II-14	281	81	15	37	414	524	493	6	142	7	2	4	1038	87	9	67	256	3463	0.43	8069
HH-II-17	100	35	1	7	220	152	173	1	97	1	1	4	504	36	7	16	108	1463	0.53	2786
HH-III	42	6	0	2	93	67	65	0	21	1	2	0	96	57	3	8	64	527	0.55	952
HH-III-3	96	32	8	18	625	305	245	0	74	6	3	0	310	72	5	74	291	2164	0.64	3397
HH-III-4	295	12	1	21	192	128	98	0	36	3	5	0	204	63	3	23	229	1313	0.76	1725
HH-III-5	144	38	16	35	301	205	231	0	34	2	2	0	388	76	11	32	219	1734	0.48	3612
HH-III-6	134	29	5	18	290	183	197	9	58	2	3	0	426	66	10	33	180	1643	0.46	3540
HH-III-8	92	26	4	6	528	203	260	0	722	1	2	0	294	154	4	29	255	2580	0.68	3774
HH-III-9	109	22	0	12	239	143	196	0	87	3	4	0	462	63	7	38	164	1549	0.49	3918
HH-III-12	658	97	40	32	369	318	217	0	98	3	6	0	1071	135	20	121	477	3662	1.03	3540
HH-III-13	119	29	8	14	226	176	246	3	53	6	2	0	345	70	6	28	142	1473	0.41	3576
HH-III-14	133	33	10	11	242	184	195	1	65	2	21	0	408	86	5	39	146	1581	0.45	3504
HH-III-17	49	6	0	3	123	81	86	0	23	2	2	0	154	67	4	11	74	685	0.49	1402
GenBank relative size	0.14	0.03	0.01	0.01	0.07	0.07	0.07	<0.01	0.03	<0.01	<0.01	<0.01	0.37	0.02	0.01	0.06	0.10	1.00		

The motifs are named as explained in Methods and in Figure 1. The “expected” number of occurrences was obtained by searching the different motifs in a database of 1000 random sequences of 100,000 nucleotides (equal representations of A, C, G, and T) and correcting the frequency to the relative size of the GenBank. The “total over expected” shows the ratio of occurrences obtained in the GenBank versus the expected ones according to the search performed in a random database with the size of the GenBank.

(pri) Primate sequence entries (from the two GenBank files); (rod) rodent sequence entries; (mam) other mammalian sequence entries; (vrt) other vertebrate sequence entries; (inv) invertebrate sequence entries; (pln) plant sequence entries (including fungi and algae); (bct) bacterial sequence entries; (rna) structural RNA sequence entries; (vrl) viral sequence entries; (phg) phage sequence entries; (syn) synthetic and chimeric sequence entries; (una) unannotated sequence entries; (est) EST (expressed sequence tag) (from 23 GenBank files); (pat) patent sequence entries; (sts) STS (sequence tagged site) sequence entries; (gss) GSS (genome survey sequence) sequence entries; (htg) HTGS (high throughput genomic sequencing) sequence entries.

Table 3. Distribution of Hammerhead and Hammerhead-like Motifs in the Different Sections of the GenBank: Mutants of Helices II and III

	pri	rod	mam	vrt	inv	plh	bct	rna	vrl	phg	syn	una	est	pat	sts	gss	htg	Total	Total over expected	Expected (A = C = G = T)
HH-I	51	21	0	14	176	119	130	0	37	3	5	0	186	45	4	17	108	916	0.56	1635
HH-I-iIAU	569	100	30	36	822	545	426	3	58	7	8	0	1113	93	32	341	544	4727	2.56	1851
HH-I-iICG	343	59	12	20	203	117	512	11	24	2	1	11	669	69	15	66	221	2355	1.27	1851
HH-I-iIGU	452	65	9	38	361	227	203	0	73	3	49	2	794	58	15	93	332	2774	1.77	1564
HH-I-iIUA	606	124	21	82	406	364	260	2	165	10	2	0	941	43	15	175	436	3652	2.03	1797
HH-I-iIUG	455	69	25	38	354	268	287	4	57	6	24	2	664	47	9	92	268	2669	1.43	1869
HH-I-iIiCG	51	19	5	6	91	63	163	0	43	2	2	0	355	77	1	10	27	915	0.48	1887
HH-I-iIiCC	121	53	13	13	103	116	189	1	48	1	6	0	494	60	2	28	52	1300	0.74	1761
HH-I-iIiCGU	77	31	18	12	98	116	130	0	21	0	0	1	284	44	1	17	64	914	0.58	1564
HH-I-iIiUA	52	23	2	8	118	89	79	2	13	0	8	0	177	28	2	18	38	657	0.39	1707
HH-I-iIiUG	36	19	4	6	100	63	155	6	21	2	152	0	154	73	1	19	90	901	0.50	1797
HH-II	83	33	1	5	175	126	141	1	80	1	1	3	435	30	4	8	91	1218	0.54	2246
HH-II-iIAU	719	117	40	39	860	572	453	0	118	13	2	0	1021	87	24	190	663	4918	2.76	1779
HH-II-iICG	404	97	21	25	183	113	148	1	50	1	7	0	856	38	20	83	284	2331	1.34	1743
HH-II-iIGU	560	96	36	39	352	270	237	1	192	5	1	0	783	35	15	124	367	3113	1.77	1761
HH-II-iIUA	668	126	133	39	465	382	308	4	85	10	22	1	843	66	21	248	462	3883	2.12	1833
HH-II-iIUG	694	123	36	24	335	239	260	1	71	0	5	0	749	67	17	212	319	3152	1.81	1743
HH-II-iIiCG	65	28	3	14	96	45	215	2	15	4	1	0	159	10	1	20	49	727	0.40	1833
HH-II-iIiCC	136	67	16	21	92	103	232	0	20	7	1	2	498	51	2	19	72	1339	0.71	1887
HH-II-iIiCGU	69	42	3	12	101	88	182	0	33	6	2	0	365	30	1	19	65	1018	0.48	2103
HH-II-iIiUA	60	16	3	11	107	78	102	1	27	0	0	0	136	8	1	21	65	636	0.35	1833
HH-II-iIiUG	50	22	4	27	85	77	129	0	39	0	0	0	154	23	0	17	63	690	0.30	2282
HH-III	42	6	0	2	93	67	65	0	21	1	2	0	96	57	3	8	64	527	0.55	952
HH-III-iIAU	360	59	17	11	482	324	241	0	35	7	11	1	723	31	6	354	328	2990	5.05	593
HH-III-iICG	203	37	4	9	98	58	101	0	9	0	0	0	469	8	6	42	132	1176	1.17	1006
HH-III-iIGU	251	29	3	19	194	126	121	0	43	4	0	0	440	40	13	52	217	1552	1.73	899
HH-III-iIUA	333	48	10	23	246	224	153	13	39	1	0	0	486	22	5	70	257	1930	2.24	863
HH-III-iIUG	285	56	11	31	212	139	113	0	29	0	1	0	399	27	8	63	148	1522	2.02	755
HH-III-iIiCG	23	9	2	3	45	22	108	0	8	1	0	0	158	52	1	7	19	458	0.50	917
HH-III-iIiCC	78	14	24	15	38	66	100	0	12	2	1	0	173	27	2	18	38	608	0.57	1060
HH-III-iIiCGU	52	18	10	13	47	51	83	1	18	1	11	1	235	7	4	9	43	604	0.53	1150
HH-III-iIiUA	43	15	7	3	72	64	137	2	8	3	304	0	98	133	3	18	126	1036	1.56	665
HH-III-iIiUG	26	17	6	5	40	33	82	3	17	1	152	0	97	64	4	8	58	613	0.67	917
GenBank relative size	0.14	0.03	0.01	0.01	0.07	0.07	0.07	<0.01	0.03	<0.01	<0.01	<0.01	0.37	0.02	0.01	0.06	0.10	1.00		

See legend of Table 2 for explanation of abbreviations.

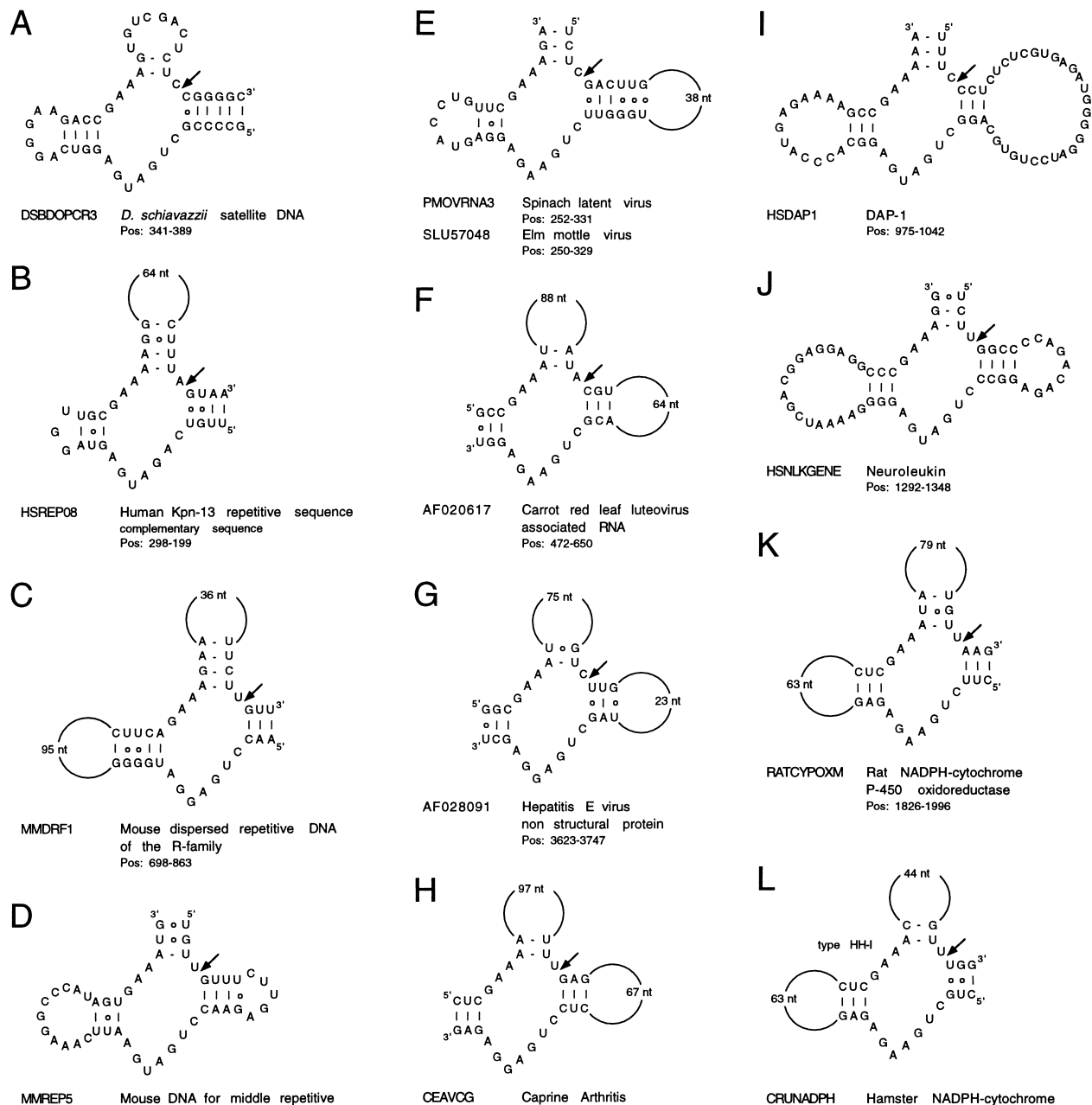


Figure 2 A–L show putative hammerhead motifs.

disabling A at position 4, but one (AA564135) possesses a C. The latter motif is inactivated by a G per A substitution at position 12. Variants of this motif also are found in genomic clones containing Kpn repetitive sequences. Intriguingly, the L1 motif interrupting the dystrophin gene of a muscular dystrophy patient (accession number HSU09115) also has a disruption in Helix I. Four additional hammerhead-like motifs were found in the satellite DNA array from the rodent *Microtus chrotorrhinus* (accession number MICSATB, posi-

tion 921–1079, not shown), in the repetitive DNA from the protozoan parasite *Theileria parva* (accession number S37077, position 84–223, not shown) with the descriptor HH-I-7 and in mouse repetitive DNA with descriptors for the HH-I-iiUA and HH-III-iiAU motifs (Fig. 2C,D). The first two motifs are predicted to be inactive because they contain A instead of G in position 12.

Viruses are good candidates for using catalytic RNA motifs. We have found several new intriguing hammerhead motifs in different viruses (Fig. 2E). Two

similar hammerhead ribozyme motifs were found in the 5' untranslated region of two viruses of the *Ikarvirus* genus, family of Bromoviridae, which are single stranded positive RNA viruses. One motif is in the spinach latent virus (accession number PMOVRNA3, position 252–331) and the other in the Elm mottle virus (accession number SLU57048, position 250–329) (Fig. 2E). Both motifs were found using the HH-III descriptor. The region containing the hammerhead is highly conserved among these viruses. The hammerhead motif found with HH-II in an RNA associated to carrot red luteovirus that is also very interesting because satellite RNAs were the first molecules found to contain hammerhead ribozymes (Fig. 2F). This motif is predicted to cleave after AUA. Mammalian viruses also contain potential hammerhead ribozymes, and two of them found with HH-II are illustrated in Figure 2G,H, one in the hepatitis E virus, and the other in the caprine encephalitis virus.

Two hammerhead motifs in human mRNAs also are presented in Figure 2I,J. Self-cleaving motifs in mRNA might regulate gene expression by promoting RNA decay. The genes coding for the interferon-induced DAPI and the neuroleukin gene possess potentially active hammerhead motifs found with HH-III that are predicted to cleave after UUC and CUC, respectively. Perhaps even more remarkable are the conserved hammerhead motifs found in the genes coding for NADPH-cytochrome P450 oxidoreductase both in the rat and the hamster (Fig. 2K,L). All together, the motifs presented here suggest that the hammerhead ribozyme might have functions other than those previously suggested for satellite RNA and transcripts for repetitive sequences.

DISCUSSION

We have used the search engine RNAMOT to scrutinize the GenBank for potential self-cleaving hammerhead ribozyme motifs. Our search extends earlier efforts to find a subset of potential hammerheads in *Escherichia coli* sequences (Ruffner et al. 1990). Because this motif has relatively few structural constraints, we designed an extensive set of descriptors for both the wild-type motif and variants of its essential nucleotides. The results show a wide distribution of potential hammerhead-like motifs in all regions of the GenBank with a higher frequency for the variants that do not require the presence of a CpG dinucleotide in the final sequence of the motifs. This CpG dinucleotide in positions 11.1 and 12 is not absolutely required for self-cleavage because other base pairs are acceptable in positions 10.1:11.1. We conclude that the reduction we observed in the frequency of most hammerhead motifs in this search is fortuitous.

We expect that most of the motifs found here are inactive because we designed descriptors that include

mutations or nonoptimal features of the hammerhead self-cleaving motif (Ruffner et al. 1990). However, our results illustrate the possibility that natural sequences might end up forming self-cleaving motifs by random drift. In other words, it would be sufficient to mutate one or two residues to activate the potential hammerhead ribozymes described here. This is not only true for the hammerhead ribozyme motif because other RNA motifs can be found randomly in natural sequences (Fontana et al. 1993; Reidys et al. 1997; Bourdeau et al. 1999).

The use of variants of the hammerhead ribozyme was stimulated by previous work that showed that satellite DNA encoding hammerhead ribozymes is enriched with mutated variants of the motif (Zhang and Epstein 1996; Ferbeyre et al. 1998). The ribozyme motif found in the cricket satellite DNA follows this rule because 14 of the 20 sequences deposited until now in the GenBank contains an active motif. Other mutant hammerheads were found in different families of repetitive DNA by using descriptors for hammerhead-like motifs, raising the possibility that other members of these families, not yet sequenced, contain the active motifs. The occurrence of hammerhead ribozymes in transcripts of repetitive DNA from different species suggests a functional role for the self-cleavage reaction in the propagation and/or the metabolism of these transcripts. We previously have proposed that self-cleavage might limit the expansion of repetitive sequences through the genome by retrotransposition (Ferbeyre et al. 1998). This model predicts that recent insertions of these elements will contain disabling mutations in the hammerhead motif. The family of L1 repetitive elements for example contains mutated versions of the hammerhead and members of this family still retrotranspose in humans, sometimes causing genetic diseases (Holmes et al. 1994). Another intriguing possibility is that viroids and satellite RNAs originated from transcripts of repetitive sequences when these transcripts parasitizes a viral replication machinery. Subsequently, they might jump from one organism to another using the virus as a vector, and as a result their distribution will cross phylogenetic barriers.

Many ESTs and mRNAs were found here to possess hammerhead-like motifs. To test any role of the hammerhead motifs identified in this work, we need a combination of biochemical and genetic analysis. Our group has finished the characterization of hammerhead motifs in repetitive DNA of Schistosome (Ferbeyre et al. 1998) and the cricket (A.A. Rojas, A. Vazques-Tello, G. Ferbeyre, F. Venanzetti, L. Bachmann, B. Paquin, and R. Cedergren, in prep.). All the occurrences we found in the GenBank are available at our web site (URL: <http://www.centrcn.umontreal.ca/~bourdeav/HH>) for those interested in finding where "hammers" can cut.

METHODS

The pattern searching for RNA secondary structures was performed by RNAMOT (Gautheret et al. 1990; Laferrière et al. 1994). The inputs for this program are nucleotide sequences, and a descriptor file defining the structural motif to be searched. RNAMOT reports all the occurrences of the motif as well as its positions along the sequence. Two of the three helices defining the hammerhead self-cleaving motif are closed by loops. The remaining helix connects the motif to the rest of the RNA molecule. As a result, there are three ways of defining a self-cleaving hammerhead ribozyme motif. We have built descriptors for these three different orientations of the motif taking into account the following constraints (Fig. 1):

1. Three nucleotides in Helix I. Helix I has no specific nucleotide requirements although the hammerhead motif found in the newt and in Schistosoma possess a conserved GG:CC base pairing, three nucleotides downstream from the cleavage site as well as an internal loop farther downstream (Pabon-Peña et al. 1991; Ferbeyre et al. 1998).
2. The conserved sequence CYGANGA. This sequence is part of the catalytic core of the ribozyme and is entirely conserved with the exception of position 7. In the latter, although all nucleotides are accepted, the preferred ones are U then G or A and finally C. More recently, position 4 was reported to accept also U, so we have included this feature in our search (Ambros and Flores 1998).
3. Three nucleotides in Helix II. There is a strong preference for a R:Y base pair in positions 10.1:11.1, but the pair G:C confers the better activity and was the only one allowed in our original descriptors.
4. The conserved sequence GAA is absolutely required for catalysis. In the X-ray model of the hammerhead, nucleotides G12 and A13 form two reverse Hoogsteen G-A base pairs with nucleotides A9 and G8, respectively, whereas A14 form a non-Watson Crick base pair with N7 (Scott et al. 1995).
5. Helix III requires an A:U base pair which is also of non Watson Crick type and a minimum of one more pair in two of the orientations (HH-I and HH-II). When the helix is open as in HH-III, two more pairs are required.
6. The cleavage site was defined as NUH (H is any nucleotide but G). However, natural ribozymes contain GUC, GUA, AUA, and AUC because they allow the highest reaction rates (Shimayama et al. 1995; Ferbeyre et al. 1998).
7. The loops closing the helices were allowed to have from 0 to 100 nucleotides.

Sixty-three additional mutants also were included in the study. These were derived from the original motifs shown in Figure 1 by changing either one base in the conserved single stranded regions for an N (any nucleotide; 30 mutants), the identity of one of the constrained base pair (positions 10.1:11.1 and 15.1:16.1; 30 mutants), or by changing the cleavage site from NUH to NHH (three more motifs; Kore et al. 1998).

The search was performed in the July 15, 1998 release of the GenBank sequence database (National Center for Biotechnology Information-GenBank flat file release 108.0). Searches were performed on both strands and all occurrences of motifs involving unidentified bases denoted by N in the database were disregarded. A Power Challenge XL with 32 CPUs IP 19, R4400, 150-MHz processor (3072 Mbytes) running UNIX IRIX 6.2 was used.

To help establish the significance of their presence, fre-

quencies of each motif in the database were compared with frequencies in a random sequence database generated by a uniform pseudo-random number generator (L'Écuyer and Andres 1997) with a period length near 2121. The random sequence databases contained 1000 sequences of 100,000 nucleotides each; the four nucleotides A, C, G, and T were used with equal probabilities. An "expected" frequency N in GenBank was calculated from the number M of occurrences of each motif in the random databases as follows: $N = (a \times M) / (10^4 \times 10^5)$, where a is the number of nucleotides in GenBank (1.797×10^9 in the release 108.0).

The random database reduced in CpG dinucleotides was generated using the same procedure, but each time a CpG dinucleotide was created a second generator (evolving in parallel) would enter in function to decide if yes or no (50% frequency) the dinucleotide would be changed. If a change had to take place, a third generator (also evolving in parallel) would be able to choose among six replacing dinucleotides: AG, CA, CC, CT, GG, or TG (choices made according to the dinucleotide frequencies reported by Karlin and Mrazek 1997). The expected frequency was evaluated as before.

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REFERENCES

- Ambros, S. and Flores, R. 1998. *In vitro* and *in vivo* self-cleavage of a viroid RNA with a mutation in the hammerhead catalytic pocket. *Nucleic Acids Res.* **26**: 1877-1883.
- Benson, D.A., Boguski, M.S., Lipman, D.J., Ostell, J., Ouellette, B.F., Rapp, B.A., and Wheeler, D.L. 1999. GenBank. *Nucleic Acids Res.* **27**: 12-17.
- Birikh, K.R., Heaton, P.A., and Eckstein, F. 1997. The structure, function and application of the hammerhead ribozyme. *Eur. J. Biochem.* **245**: 1-16.
- Bourdeau, V., Ferbeyre, G., Pageau, M., Paquin, B., and Cedergren, R. 1999. The distribution of RNA motifs in natural sequences. *Nucleic Acids Res.* **27**: 4457-4467.
- Bratty, J., Chartrand, P., Ferbeyre, G., and Cedergren, R. 1993. The hammerhead RNA domain, a model ribozyme. *Biochim. Biophys. Acta* **1216**: 345-359.
- Bussi re, F., Lafontaine, D., and Perreault, J.-P. 1996. Compilation and analysis of viroid and viroid-like RNA sequences. *Nucleic Acids Res.* **24**: 1793-1798.
- Epstein, L.M. and Gall, J.G. 1987. Self-cleaving transcripts of satellite DNA from the newt. *Cell* **48**: 535-543.
- Ferbeyre, G., Smith, J.M., and Cedergren, R. 1998. Schistosoma satellite DNA encodes active hammerhead ribozymes. *Mol. Cell. Biol.* **18**: 3880-3888.
- Fontana, W., Konings, D.A., Stadler, P.F., and Schuster, P. 1993.

- Statistics of RNA secondary structures. *Biopolymers* **33**: 1389–1404.
- Forster, A.C. and Symons, R.H. 1987. Self-cleavage of plus and minus RNAs of a virusoid and a structural model for the active sites. *Cell* **49**: 211–220.
- Gautheret, D., Major, F., and Cedergren, R. 1990. Pattern searching/alignment with RNA primary and secondary structures: an effective descriptor for tRNA. *Comput. Appl. Biosci.* **6**: 325–331.
- Hertel, K.J., Pardi, A., Uhlenbeck, O.C., Koizumi, M., Ohtsuka, E., Uesugi, S., Cedergren, R., Eckstein, F., Gerlach, W.L., and Hodgson, R., et al. 1992. Numbering system for the hammerhead. *Nucleic Acids Res.* **20**: 3252.
- Holmes S.E., Dombroski B.A., Krebs C.M., Boehm, C.D., and Kazazian, H.H. Jr. 1994. A new retrotransposable human L1 element from the LRE2 locus on chromosome 1q produces a chimaeric insertion. *Nat. Genet.* **7**: 143–148.
- Karlin, S. and Mrazek, J. 1997. Compositional differences within and between eukaryotic genomes. *Proc. Natl. Acad. Sci.* **94**: 10227–10232.
- Kore, A.R., Vaish, N.K., Kutzke, U., and Eckstein, F. 1998. Sequence specificity of the hammerhead ribozyme revisited; the NHH rule. *Nucleic Acids Res.* **26**: 4116–4120.
- Laferrère, A., Gautheret, D., and Cedergren, R. 1994. An RNA pattern matching program with enhanced performance and portability. *Comput. Appl. Biosci.* **10**: 211–212.
- Lafontaine, D.A., Deschènes, P., Bussière, F., Poisson, V., and Perreault, J.-P. 1999. The viroid and viroid-like RNA database. *Nucleic Acids Res.* **27**: 186–187.
- L'Écuyer, P. and Andres, T.H. 1997. A random number generator based on the combination of four LCGs. *Math. Comput. Simulation* **44**: 99–107.
- Pabon-Peña, L.M., Zhang, Y., and Epstein, L.M. 1991. Newt satellite 2 transcripts self-cleave by using an extended hammerhead structure. *Mol. Cell. Biol.* **11**: 6109–6115.
- Reidys, C., Stadler, P.F., and Schuster, P. 1997. Generic properties of combinatorial maps: Neutral networks of RNA secondary structures. *Bull. Math. Biol.* **59**: 339–397.
- Ruffner, D.E., Stormo, G.D., and Uhlenbeck, O.C. 1990. Sequence requirements of the hammerhead RNA self-cleavage reaction. *Biochemistry* **29**: 10695–10702.
- Scott, W.G., Finch, J.T., and Klug, A. 1995. The crystal structure of an all-RNA hammerhead ribozyme: A proposed mechanism for RNA catalytic cleavage. *Cell* **81**: 991–1002.
- Shimayama, T., Nishikawa, S., and Taira, K. 1995. Generality of the NUX rule: Kinetic analysis of the results of systematic mutations in the trinucleotide at the cleavage site of hammerhead ribozymes. *Biochemistry* **34**: 3649–3654.
- Symons, R. 1992. Small catalytic RNAs. *Annu. Rev. Biochem.* **61**: 641–671.
- Symons, R. 1997. Plant pathogenic RNAs and RNA catalysis. *Nucleic Acid Res.* **25**: 2683–2689.
- Zhang, Y. and Epstein, L.M. 1996. Cloning and characterization of extended hammerheads from a diverse set of caudate amphibians. *Gene* **172**: 183–190.

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