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COMMENTARY

A Concern Regarding the Current Confusion with the Human Homolog of Mouse Np95, ICBP90/UHRF1

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ICBP90/UHRF1, which is overexpressed in cancer cells and is down-regulated by p53, possesses a methylated CpG binding affinity and binds to the methylated promoters of tumor suppressor genes in cancer cells with HDAC1 and DNMT1, suggesting suppression of these genes and maintenance of methylation status which leads to carcinogenesis. Recently, it was reported that the human homolog of Np95 is different from ICBP90 but not from UHRF1. Because UHRF1 is the gene symbol of ICBP90, the claim is a little confusing; that is, UHRF1 and ICBP90 are identical. Because the previously published genomic structure of the ICBP90 gene needed to be revised and the registered ICBP90 sequence (AF129507) contains two rare polymorphisms or sequence errors, we think that confusion could occur. Here we show the revised ICBP90 gene structure and 366 polymorphisms in this gene. Our conclusion is that the human homolog of Np95 is ICBP90, whose gene symbol is UHRF1. © 2008 by Radiation Research Society

INTRODUCTION

ICBP90 (gene symbol *UHRF1*), which was first identified as a *topoisomerase II alpha* regulator, is a very important protein that is associated with cell cycle progression and is overexpressed in proliferating cells and cancer cells (1-3). ICBP90 has a binding affinity to methylated CpGs, makes a complex with HDAC1, and binds to the promoters of methylated tumor suppressor genes, suggesting that ICBP90 promotes cancer cell proliferation through suppression of these tumor suppressor genes (2). ICBP90 is also down-regulated by p53 (4), possesses ubiquitin E3 li-

¹ Address for correspondence: Laboratory of Human Carcinogenesis, Center for Cancer Research, National Cancer Institute, NIH, 37 Convent Drive, Building 37, Room 3060, Bethesda, Maryland 20892-4258; e-mail: unokim@mail.nih.gov. gase activity (5), and plays a role in maintaining DNA methylation with DNMT1, keeping the genes silent (6, 7). ICBP90 has three family proteins, NIRF/UHRF2, ICBP55/UHRF3 and ICBP87/UHRF4, that share 52.6, 59.8 and 78.1% identity with ICBP90, respectively (3).

Recently an article titled "Isolation and Characterization of a Novel Human Radiosusceptibility Gene, *NP95*" was published in *Radiation Research* (8). The authors claimed that they identified a new human homolog of mouse Np95 and the "new" gene was identical to *UHRF1* but not to *ICBP90*. However, because the gene symbol of ICBP90 is *UHRF1* (i.e., they are the same) and the *ICBP90* sequence has been registered for a long time (AF129507, submitted by Hopfner *et al.* on Feb. 19, 1999), we are afraid that the article will create major confusion in the future. We would like to discuss the possibilities that could have led to the confusion.

MATERIALS AND METHODS

Databases

We used the databases below to determine the genomic DNA structure of *ICBP90* and polymorphisms in the gene: BLAST (http://www.ncbi. nlm.nih.gov/BLAST/), Unigene (http://www.ncbi.nlm.nih.gov/sites/ entrez?db=unigene), Ensemble (http://www.ensembl.org/index.html), GenBank (http://www.ncbi.nlm.nih.gov/Genbank/), DDBJ (http://www. ddbj.nig.ac.jp/), dbSNP (http://www.ncbi.nlm.nih.gov/sites/entrez?db= snp), and JSNP (http://snp.ims.u-tokyo.ac.jp/).

RESULTS AND DISCUSSION

Concerning the recently published article entitled "Isolation and Characterization of a Novel Human Radiosusceptibility Gene, NP95" (8), we have a strong concern that the authors did not isolate a new gene. In the discussion of the article, it is described that they could not recognize the sequence of exon F (2.6 kb) of the *ICBP90* gene in the *Homo sapiens* chromosome 19 genomic contig, NT_011255 (GenBank), at 19p13.3, but found this exon in a BAC

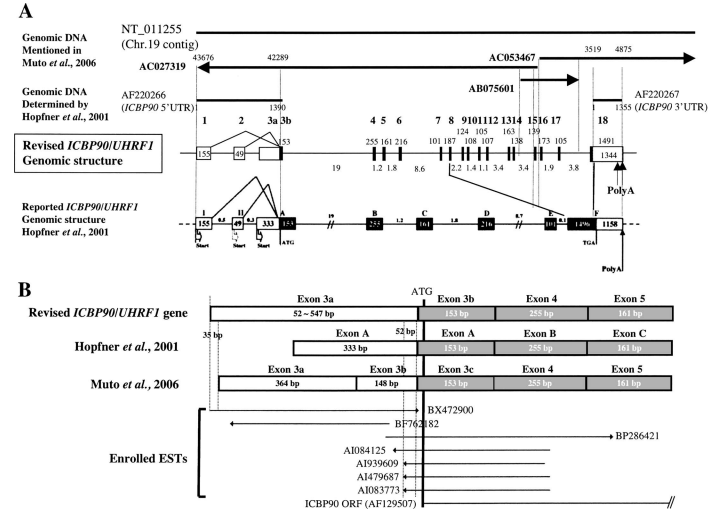


FIG. 1. Panel A: Revised full-length *ICBP90/UHRF1* gene on chromosome 19 contig, NT_01125. AC027319, AC053467 and AB075601 are the three genomic DNA fragments that were mentioned by Muto *et al.* (8). Exon F reported by Hopfner *et al.* (9) was composed of 11 exons. AF220266 and AF220267 were the *ICBP90* 5'UTR and 3'UTR sequence determined by Hopfner *et al.* (9). Panel B: Exon 3 is composed of exons 3a and 3b. The length of exon 3a is 52–547 bp based on the seven registered ESTs. Exon 3b is 153 bp including the start codon (ATG) and is involved in all transcripts.

clone, AC112777, that includes 12p12.2-p12.1. Hopfner et al. (9) have reported that the ICBP90 gene spans approximately 35.8 kb on chromosome 19p13.3 and contains six coding exons named A to F. We carefully checked the genomic structure of the ICBP90 gene and found that exon F is actually composed of 11 exons (Fig. 1). All exons of ICBP90 are on the contig, NT_011255, which is composed of many genomic fragments, including AC027319, AC053467 and AB07561, which are discussed in the article. Exons 1 to 14 are on AC027319, exons 15 to 17 are on AB075601, and exons 16 to 18 are on AC053467. Therefore, the ICBP90 gene is localized at the same position where the "new" human NP95 homolog is located. We think that this incorrect definition of exon F (9) is one of the major sources of confusion that led Muto et al. to think that the ICBP90 gene is different from the gene that they identified.

It is also stated in their discussion that human NP95 and

UHRF1 are identical in ORF and that *ICBP90* and human *NP95* differ in ORF by only two amino acids (Lys383 \rightarrow Asn383, Ala457 \rightarrow Ser457) and seven nucleotides. A difference of nine nucleotides in a long identical sequence (in this case, *ICBP90*'s ORF is 2382 bp) does not make it a different gene. Usually these differences are polymorphisms or sequence errors. Therefore, we searched SNP databases and found 366 polymorphisms including 308 single nucleotide polymorphisms (SNPs) and 58 deletions on the gene (Table 1, Supplemental Table 1). Among these 366 polymorphisms, 337 were located in introns, six were in 3'UTR, one was in 5'UTR, and 22 were in exons (Table 1). Among the 22 polymorphisms in exons, 15 were synonymous substitutions and seven were nonsynonymous substitutions (Table 2).

We are not sure whether all nine nucleotide differences are polymorphisms, because there are no descriptions of seven of the nine different nucleotides in the article. How-

Polymorphisms in the <i>ICBP90/UHRF1</i> Gene							
Location	Deletion	Substitution (SNP)	Synonymous substitutions	Nonsynonymous substitutions	Total polymorphism numbers		
Intron	58	279	0	0	337		
3'UTR	0	6	0	0	6		
5'UTR	0	1	0	0	1		
Coding region	0	22	15	7	22		
Total	58	308	15	7	366		

 TABLE 1

 Polymorphisms in the ICBP90/UHRF1 Gene

ever, the two differences (Lys383 \rightarrow Asn383, Ala457 \rightarrow Ser457) that are described in the article were not involved in the 366 polymorphisms. We determined the sequence by sequencing genomic DNA from several cancer cell lines and Japanese individuals and determined that codon 383 is AAG (Lys) and codon 457 is GCG (Ala); on the other hand, codon 383 is AAT (Asn) and codon 457 is TCG (Ser) in the registered ICBP90 sequence (AF129507). Therefore, AF129507 includes at least two rare polymorphisms that are not registered in the database or are sequence errors at codons 383 and 457. The other seven different nucleotides could be polymorphisms. Unoki et al. (2) used ICBP90 plasmid vectors that include Lys383 and Ala457, not Asn383 and Ser457, and three different polymorphisms compared with AF129507, because these polymorphisms were major alleles in the database. If the claim of Muto et al. (8) is right, Unoki et al. (2) did not use ICBP90 in their experiments. But no one accepts that two genes at exactly

the same location in a chromosome with 99.748% amino acid identity (only two amino acids are different among 754 amino acids) are different.

Our other concern is the 5'UTR splice variations that are described in Fig. 2 (8). Hopfner *et al.* (9) reported three 5'UTR variations that are different from the variations in their article. We checked the EST database using BLAST and found that the three different 5'UTR variations that Hopfner *et al.* (9) described were registered as EST fragments, indicating that they are actually expressed in cells (Fig. 2, Supplemental Table 2). However, there are no ESTs that include exon 1 and 2 as described by Muto *et al.* (8). Transcripts that include these exons may be expressed at a low level, but we need expression data. At the least, they should add the 5'UTR variation that Hopfner *et al.* (9) identified in their article.

The authors also claimed that their newly isolated gene differs in its effect on topoisomerase II activity. However,

dbSNP rs number		Flanking sequence	Exon location	
15 Syne	onymous SNPs			
1	rs2261986	TGTGGATCCAGGTTCGGACCATGGA [C/T] GGGAGGCAGACCCACACGGTGGACTC	exon 3	
2	rs2251520	GAGTCAGACAAGTCCTCCACCCACGG [T/C] GAGGCGGCCGCCGAGACTGACAGCA	exon 4	
3	rs2123731	GATGAGGACATGTGGGATGAGACGGA [A/G] TTGGGGCTGTACAAGGTGAGCCTCC	exon 4	
4	rs2307205	TCCCGGGACGAGCCCTGCAGCTCCAC [G/A] TCCAGGCCGGCGCTGGAGGAGGACG	exon 5	
5	rs2307201	GGCGTGGTCCAGATGAACTCCAGGGA [C/T] GTCCGAGCGCGCGCCCGCACCATCA	exon 6	
6	rs2307206	CAGATGAACTCCAGGGACGTCCGAGC [G/A] CGCGCCCGCACCATCATCAAGTGGC	exon 6	
7	rs17881281	CTGGAGGTGGGCCAGGTGGTCATGCT [C/T] AACTACAACCCCGACAACCCCAAGG	exon 6	
8	rs4807002	GCCCCGTGCCCAGGGAAGAGCGGGGCC[G/A]TCCTGCAAGCACTGCAAGGACGACG	exon 8	
9	rs17883422	GACCCGCCCTCAGCAGTGTTCCCAG [C/T] GAGGACGAGTGGTGAGTGCGGCCCT	exon 8	
10	rs2250982	GTCCATCGGCCCCACGTGGCTGGCAT [C/A] CACGGCCGGAGCAACGACGGAGCGT	exon 11	
11	rs2250981	CATCGGCCCCACGTGGCTGGCATACA [C/T] GGCCGGAGCAACGACGGAGCGTACT	exon 11	
12	rs2250978	GTCCTGGCGGGGGGGCTATGAGGATGA [C/T] GTGGTGAGTGTGTGTGGGGAGGGG	exon 11	
13	rs2307213	TACGCCCCCGCTGAGGGCAACCGCTA [C/T] GATGGCATCTACAAGGTGAGTGCCC	exon 13	
14	rs17878253	AGGGAGGAGGAGGAGCAGCAGGAGGG [G/T] GGCTTCGCGTCCCCCAGGACGGGCA	exon 15	
15	rs2247238	GAGGAGGAGCAGCAGGAGGGGGGCTT [C/T] GCGTCCCCCAGGACGGGCAAGGGCA	exon 15	
7 Nons	ynonymous SNPs			
1	rs17886098	ACCCCAAGGAGCGGGGCTTCTGGTAC [G/C] ACGCGGAGATCTCCAGGAAGCGCGA	exon 6	
2	rs2292148	GCCGTCCTGCAAGCACTGCAAGGAC [A/G] ACGTGAACAGACTCTGCCGGGTCTGC	exon 8	
3	rs17885791	TGGTACTGGCGGGAGAGCGGCTGAGA [G/A] AGAGCAAGAAGAAGGCGAAGATGGC	exon 9	
4	rs2307211	CAAGTACGCCCCCGCTGAGGGCAACC [G/T] CTATGATGGCATCTACAAGGTGAGT	exon 13	
5	rs17883331	AGGAGGAGCAGCAGGAGGGGGGGCTTC [G/A] CGTCCCCCAGGACGGGCAAGGGCAA	exon 15	
6	rs17884843	GGAGGGGGGCTTCGCGTCCCCCAGGA [C/T] GGGCAAGGGCAAGTGGAAGCGGAAG	exon 15	
7	rs17883563	ACACCCGCTTCCCTCTAGTTCCAGTT [G/T] TTCCTGAGTAAAGTGGAGGAGACGT	exon 17	

 TABLE 2

 Detailed SNP Information in ICBP90/UHRF1 Gene Exons

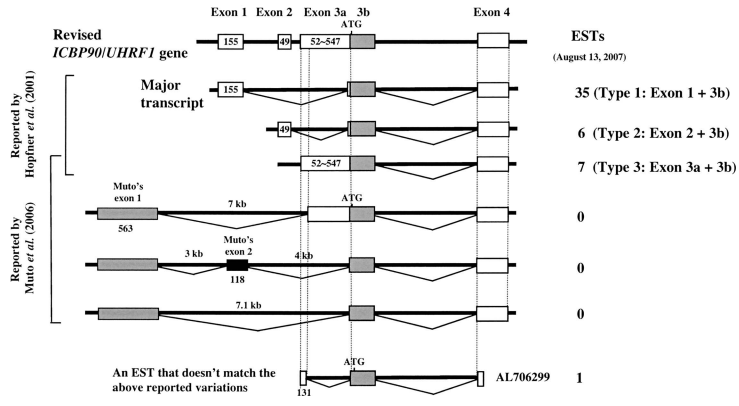


FIG. 2. 5'UTR variations of *ICBP90/UHRF1* transcripts. Major transcript (35 ESTs have been enrolled) has exon 1 and exon 3b. A transcript that has exon 2 and exon 3b (six ESTs have been registered) and another transcript that has exon 3a and exon 3b (seven ESTs have been registered) are secondarily abundant transcripts based on an EST database search. There are no ESTs corresponding to the three transcripts that are described by Muto *et al.* (8). AL706299 has a 5' part of exon 3a and exon 3b and does not match any reported transcripts.

Extended								
	Amino acid		Allele frequency					
Substitution type	Major	Minor	Major	Minor	Amino acid position			
synonymous	C: Asp	T: Asp	C: 0.7	T: 0.3	9			
synonymous	T: Gly	C: Gly	T: 0.674	C: 0.326	111			
synonymous	A: Glu	G: Glu	A: 0.688	G: 0.312	131			
synonymous	G: Thr	A: Thr	G: 0.989	A: 0.011	173			
synonymous	C: Asp	T: Asp	C: 0.994	T: 0.006	203			
synonymous	G: Ala	A: Ala	G: 0.982	A: 0.018	206			
synonymous	C: Leu	T: Leu	C: 0.994	T: 0.006	225			
synonymous	G: Pro	A: Pro	G: 0.993	A: 0.007	300			
synonymous	C: Ser	T: Ser	C: 0.980	T: 0.020	354			
synonymous	C: He	A: He	C: 0.591	A: 0.409	449			
synonymous	C: His	T: His	C: 0.522	T: 0.478	450			
synonymous	C: Asp	T: Asp	C: 0.925	T: 0.075	469			
synonymous	C: Tyr	T: Tyr	C: 0.639	T: 0.361	555			
synonymous	G: Gly	T: Gly	G: 0.977	T: 0.023	635			
synonymous	C: Phe	T: Phe	C: 0.771	T: 0.229	637			
nonsynonymous	G: Asp	C: His	G: 0.994	C: 0.006	240			
nonsynonymous	G: Asp	A: Asn	G: 0.988	A: 0.012	308			
nonsynonymous	G: Glu	A: Lys	G: 0.983	A: 0.017	379			
nonsynonymous	G: Arg	T: Leu	G: 0.987	T: 0.013	554			
nonsynonymous	G: Ala	A: Thr	G: 0.835	A: 0.165	638			
nonsynonymous	C: Thr	T: Met	C: 0.994	T: 0.006	642			
nonsynonymous	G: Leu	T: Phe	G: 0.994	T: 0.006	713			

TABLE 2 Extended the authors did not compare the activity of their "new" gene with ICBP90 in the same experiment. The difference between two proteins should be compared in the same experiment. The effect of ICBP90 on topoisomerase II α activation might not be as strong. In fact, Unoki *et al.* (2) got a similar result. They showed that ICBP90 has a stronger affinity for methylated CpG than for the CCAAT repeat at the topoisomerase promoter. However, as is well known, to get the maximum protein activity, special conditions are required (salt concentration, binding partner, etc.), and the condition that Unoki *et al.* (2) used might not be suitable to get the strongest activation of topoisomerase by ICBP90. Although the results of Muto *et al.* (8) are interesting, it is too early to conclude anything.

In conclusion, the genomic structure of ICBP90 that was defined previously was revised (Fig. 1), and the registered ICBP90 sequence (AF129507) includes at least two rare polymorphisms or sequence errors. All exons of the ICBP90 gene are on 19p13.3. ICBP90 (gene symbol UHRF1) is a human homolog of mouse Np95, as we have described before (3). Actually, because ICBP90 and mouse Np95 share 73.3-73.7% amino acid identity, the percentage is a little low compared to an ordinary homolog between human and mouse. However, because the human genome project was finished in 2003, we do not think that there is another human NP95 homolog on the human genome. If there is another one, it should share a higher percentage of amino acid identity with mouse Np95 than ICBP90. In addition, functional differences should be compared in the same experiment. Since the gene that the authors isolated and ICBP90 are identical, the authors' functional analysis data seem to suggest that ICBP90's function is the same as Unoki et al. reported (2) where they used the same sequence that the authors identified. We hope that this commentary can prevent future confusion.

SUPPLEMENTARY INFORMATION

Supplementary Table 1. Detailed information for all SNPs on the *ICBP90/UHRF1* gene. http://dx.doi.org/10. 1667/RR1209.1.s1

Supplementary Table 2. ESTs that include *ICBP90/UHRF1* 5'UTR variation (available August 13, 2007). http://dx.doi.org/10/1667/RR1209.1.s2

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