



Genomes were forged by massive bombardments with retroelements and retrosequences

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Abstract

Retroposition is an efficient route to move coding regions around the genome ‘in search’ of novel regulatory elements and to shotgun regulatory elements into the genome ‘in search’ of new target genes. The templates for such retrogenes are mRNAs, and for regulatory retronuons (nuon = any definable nucleic acid sequence) usually small non-mRNAs (snmRNAs). An example in support of the ‘master gene’ model for SINEs (short interspersed repetitive elements) is provided with neuronal BC1 RNA. Furthermore, an alternative explanation of LINE (long interspersed repetitive elements) involvement in the generation of SINEs is given. I will also argue that the status of transposable elements with respect to the host resembles more symbiosis than parasitiasis and that host defense is often lenient as if even to ‘tolerate or support’ retronuons. Finally the paradox of evolution’s lack of foresight and the future exaptive use of retronuons is being dealt with by referring to W.F. Doolittle’s ‘Hierarchical Approaches to Genome Evolution’.

Introduction

The first sequence of a large ribosomal RNA (Brosius et al., 1978) was not worth much for secondary structure predictions until additional rRNA sequences became available (Woese et al., 1983). Likewise, the scientific value of the entire human genome will multiply when not only another mammalian genome (mouse) but also additional primate sequences including that of our closest relative, the chimpanzee, will become available. I can hardly disagree with opinions voicing that the 3–5% protein coding regions of higher eukaryotic genomes will be boring and that the excitement lies in understanding the remainder of the total sequence. Consequently, it is highly significant to extend genomic sequencing of eukaryotic model organisms featuring compressed genomes (pufferfish or the plant *Arabidopsis thaliana*) to organisms with genome sizes that are in the ‘normal’ range for higher Eucarya. In vertebrates, it is evident already that retroelements and retrosequences – here termed retronuons where a

nuon is any definable nucleic acid sequence (Brosius & Gould, 1992, 1993) – have contributed the majority of the extra genomic mass. Retronuon integration has a wide spectrum of effects on the host genome ranging from none to dramatic and from negative (Deininger & Batzer, 1999) to positive. The potential impact of these retronuons on genomic function (Georgiev, 1984; Brosius, 1991; Brosius & Gould, 1992; Shapiro, 1992; McDonald, 1993; Makalowski, Mitchell & Labuda, 1994; Nouvel, 1994; Flavell, 1995; Wessler, Bureau & White, 1995; Brosius & Tiedge, 1996; Britten, 1996; Kidwell & Lisch, 1997; Schmid, 1998; Jurka, 1998; Tomilin, 1999; Brosius, 1999a, b, c) is gradually being accepted. The dimension of the effect depends on whether the event took place in somatic cells or the germ line.

Importantly, the effect on the host depends largely on the location of the newly integrated retronuon with respect to resident genes or regulatory elements. In comparison to the last review on the subject (Brosius, 1999b), updated tables with examples of

Table 1. Vertebrate regulatory elements or parts of coding regions generated by retroelements

Retronuon	Gene under its influence	Species	Ancestor of source nuon	Serves as	References
LTR	cDNA 7, cDNA _γ	Human	THE-1	Polyadenylation signal	Paulson et al. (1987)
LTR	Sex-limited protein (slp)	Mouse	5' LTR of C-type retrovirus (<i>imp1</i>)	Promoter	Stavenshagen and Robins (1988); Robins and Samuelson (1992); Ramakrishnan and Robins (1997)
LTR	Oncomodulin	Rat (but not mouse)	IAP	Promoter and first exon	Banville and Boie (1989)
LTR	MIPP	Mouse	IAP	Promoter	Chang-Yeh et al. (1991)
LTR	AF-3	Human	RTVL-H	Promoter	Feuchter et al. (1992)
LTR	AF-4 (CDC4L homology)	Human	RTVL-H	Promoter	Feuchter et al. (1992)
LTR	PLT	Human	RTVL-H	Polyadenylation signal	Goodchild et al. (1992)
LTR	cH-6	Human	RTVL-H	Polyadenylation signal	Mager (1989)
LTR	cH-7	Human	RTVL-H	Polyadenylation signal	Mager (1989)
LTR	cPB-3	Human	RTVL-H	Polyadenylation signal	Mager (1989)
LTR	PLA2L (phospholipase A2 homology)	Human	RTVL-H	Promoter	Feuchter-Murthy et al. (1993)
LTR	Calibindin D28K	Human	RTVL-H	Promoter	Liu and Abraham (1991)
LTR	ZNF80 zinc finger gene	Human	ERV9	Promoter	Di Cristofano et al. (1995)
LTR	Growth factor pleiotropin (PTN)	Human	HERV-E (RTVL-1)	Trophoblast-specific promoter	Schulte et al. (1996, 1998)
LTR	Leptin receptor (OBRa)	Human	HERV-K	Alternative splicing, inclusion of 67 LTR-derived aa into C-terminus of OBRa protein	Kapitonov and Jurka (1999)
LTR-IS	A1	Mouse	MuRRS	Polyadenylation signal	Baumrucker et al. (1988)
LTR-IS	A3	Mouse	MURRS	Polyadenylation signal	Baumrucker et al. (1988)
LTR	Aromatase	Chicken	Retrovirus	Promoter and 5' exon	Matsumine et al. (1991)
CR1	Lysozyme	Chicken	Retrovirus	Transcriptional silencer	Baniahmad et al. (1987)
L1	Thymidylate synthase	Mouse	LINE	Polyadenylation signal	Harendza and Johnson (1990)
L1	Insulin I gene	Rat	LINE	Transcriptional silencer	Laimins et al. (1986)
LINE	α_s 1-casein E	Goat	LINE	mRNA stability	Pérez et al. (1994)
L1	Apolipoprotein(a)	Human	LINE	Transcriptional enhancer	Yang et al. (1998)
L1	Proteasome activator PA28 β (PMSE2b)	Mouse	LINE	Promoter	Zaiß and Klotzel (1999)
HERV-E	Salivary amylase gene	Human	Retrovirus	Promoter	Samuelson et al. (1990); Emi et al. (1988); Ting et al. (1992)
HRES-1	Transaldolase	Human	Retrovirus	Part of the coding sequence	Banki et al. (1994)

Table 1. Continued

Retronuon	Gene under its influence	Species	Ancestor of source nuon	Serves as	References
The-1, IAP	Immunoglobulin heavy chain	Human Mouse	Retrotransposon	Protein sequence building blocks	Hakim et al. (1994)
LTR	Leptin	Human	MER11	Placental enhancer	Bi et al. (1997)
ALF	Annexin VI, interleukin-4, protein kinase C- β	Human	LINE-2	Potent T-cell-specific silencer	Donnelly et al. (1999)
Bov-B LINE	Bucentaur (<i>bbcmt</i>)	Ruminantia	LINE	Large part of protein coding sequence	Takahashi et al. (1998)

For a definition of the differences between retroelements (this table) and retrosequences (Tables 2–7) see text.

Not all examples are proven exaptations. Especially events that date back not much more than a few million years could only be potential exaptations (potaptations according to Brosius & Gould, 1992, 1993).

NGF-inducible cAMP-extinguishable retrovirus-like (NICER) elements have been described (Cho et al., 1990). No association with a gene under their control has been reported.

recruited/exapted (Gould & Vrba, 1982) retronuons in vertebrates are presented. Furthermore, tables with examples that cannot be defined as exaptations as of yet have been added. These may be termed potential exaptations or potaptations (Brosius & Gould, 1992, 1993). In addition, generation of SINEs (short interspersed repetitive elements) from ‘master genes’ and the involvement of LINEs (long interspersed repetitive elements) in their biogenesis is dealt with. Furthermore, measures of the host (and mobile element) are discussed that may mitigate negative effects of transposable elements on the host. Finally, I will expand on questions concerning the perseverance of reverse transcription in genomes, despite the lack of an obvious need in the cell, on parasitism versus symbiosis, the lack of foresight in evolution and on hierarchical approaches to genome evolution (Doolittle, 1989) in the light of retronuon biology.

Biogenesis of SINEs and effects of retronuons on neighboring genes

Transposable elements (TEs) include DNA transposons and elements that disperse via RNA intermediates, so-called retroposons. Retroposons can be further divided into retroelements and retrosequences. Retroelements comprise proviruses and LTR (long terminal repeat) retrotransposons as well as non-LTR retrotransposons such as LINEs. Their common denominator is the presence of an open reading frame (ORF) encoding reverse transcriptase at least in the (rarer) full-length versions. Retrosequences arise from virtually any cellular RNA that has the potential to be reverse transcribed in the cell: ribosomal, messen-

ger and small stable RNAs (Brosius & Gould, 1992; Nouvel, 1994; Brosius, 1999b). Reverse transcriptases encoded by the aforementioned retroelements play an important part in their biogenesis (Singer, 1995; Jurka, 1998).

Parts of retroelements are efficient modulators of gene expression

Often the remnants of retroelements such as lone LTRs or other domains have been recruited as regulatory elements that act on targeted genes as promoters, transcriptional enhancers/silencers, splice sites, polyadenylation signals and even as protein coding domains (Table 1).

Small RNA derived retrosequences are frequent modulators of gene expression

Retrosequences derived from small RNA templates (mainly SINEs) whose generation apparently depends on retroelement encoded reverse transcriptases play a similar role as parts of the aforementioned retroelements in regulating targeted genes (Table 2).

Models of SINE amplification: most repeats are transposable or only certain master genes can give rise to new SINEs

There are still two conflicting models concerning the mode of SINE generation as summarized by (Brookfield, 1994): “One is that they are non-autonomous transposable elements. Here the Alu sequence transposes sequentially from site to site, retaining its capacity for further transposition. . . . This model sees

Table 2. Vertebrate regulatory elements or parts of coding regions generated by retrosequences

Retronuon	Gene/nuon that is under its influence	Species	Ancestor of source nuon	Serves as	References
B2	MHC class I genes	Mouse	tRNA ^{Lys}	Polyadenylation signal	Kress et al. (1984)
B2	B2 ⁺ mRNA _x	Mouse	tRNA ^{Lys}	Polyadenylation signal	Ryskov et al. (1984)
B2	Glutathione S-transferase	Mouse	tRNA ^{Lys}	Polyadenylation signal	Rothkopf et al. (1986)
B2	Various	Rodents	rRNA ^{Lys}	mRNA stability	Clemens (1987)
B2	Fourth component of complement (C4) in H-2 ^k haplotype	Mouse	tRNA ^{Lys}	Located in intron 10; reduces expression rate to 1/10 of non-H-2 ^k mice	Zheng et al. (1992)
B2	Muscle γ -phosphorylase kinase	Mouse	tRNA ^{Lys}	Polyadenylation signal	Maichele et al. (1993)
B2	MOK-2 zinc-finger protein	Mouse	tRNA ^{Lys}	Exerts a negative <i>cis</i> -acting effect on <i>MOK-2</i> promoter activity	Arranz et al. (1994)
B2	Leukemia inhibitory factor receptor (LIFR)	Mouse	tRNA ^{Lys}	Generating new splice variant that leads to a soluble form of LIFR	Owczarek et al. (1996); Michel et al. (1997)
C repeats	MHC	Rabbit	tRNA ^{Gly}	Polyadenylation signal	Rebiere et al. (1987); Krane and Harsdison (1990)
C repeats	Major apoprotein of pulmonary surfactant	Rabbit	tRNA ^{Gly}	Polyadenylation signal	Boggaram et al. (1988); Krnae and Hardison (1990)
C repeats	Cytochrome P-450 isozyme 4	Rabbit	tRNA ^{Gly}	Polyadenylation signal	Okino et al. (1985); Krane and Hardison (1990)
CHR-1 repeats	EP3B and EP3C prostaglandin E2 receptors	Bovine	tRNA ^{Glu}	Protein domain	Shimamura et al. (1998)
ID		Rat	tRNA ^{Ala} via neural BC1 RNA	Enhancer	McKinnon et al. (1986)
ID	pIL2	Rat	tRNA ^{Ala} via neural BC1 RNA	mRNA stability	Glaichenhaus and Cuzin (1987)
B1	pIL2, pIL8	Mouse	SRP RNA	mRNA stability	Vidal et al. (1993)
B1	Immunoglobulin κ light chain	Mouse	SRP RNA	Negative regulation of transcription	Saksela and Baltimore (1993)
Alu	Haptoglobin related gene	Human	SRP RNA	Transcriptional enhancer	Oliviero and Monaci (1988)
Alu	θ 1 globin	Higher primates	SRP RNA	CCAAT box of promoter	Kim et al. (1989)
Alu	ε -globin	Human	SRP RNA	Transcriptional modulation	Wu et al. (1990)
Alu	7.02 bidirectional promoter	Monkey	SRP RNA	Transcriptional reducer	Saffer and Thurston (1989)
Alu	c-myc	Human	SRP RNA	Transcriptional modulation	Tomilin et al. (1990)
Alu	Adenosine deaminase	Human	SRP RNA	Transcriptional enhancer	Aronow et al. (1992)
Alu	Proliferating cell nuclear antigen (PCNA)	Human	SRP RNA	Transcriptional silencer	Sell et al. (1992)
Alu	Mitochondrial hinge protein	Human	SRP RNA	Transcriptional enhancer	Liu and Bradner (1993)
Alu	SV40 origin	Human	SRP RNA	Transcriptional enhancer	Saegusa et al. (1993)
Alu	Fc ϵ RI- γ	Human	SRP RNA	Transcriptional regulation (positive and negative)	Brini et al. (1993)

Table 2. Continued

Retronuon	Gene/nuon that is under its influence	Species	Ancestor of source nuon	Serves as	References
Alu	Keratin 18 (human)	Mouse (trans-genic)	SRP RNA	Transcriptional insulation; Alus provide retinoic acid receptor binding sites	Thorey et al. (1993); Neznanov and Oshima (1993); Vansant and Reynolds (1995)
Alu	CD8 α	Human	SRP RNA	Transcriptional enhancer (located in last intron)	Hambor et al. (1993)
Alu	α -3 acetylcholine receptor subunit	Human	SRP RNA	Alternative splicing	Mihovilovic et al. (1993)
Alu	In several protein coding regions	Primates	SRP RNA	Generating new splice variants, potentially contributing new protein domains	Reviewed in Makalowski et al. (1994)
Alu	Interferon receptor, IFNR ₁ R-2	Human	SRP RNA	Alt. splicing, part of protein cod. region	Mullersman and Pfeffer (1995)
Alu-J	Double-stranded RNA-specific editase (RED1/ADAR2)	Human	SRP RNA	40 Alu-derived aa are added via alternative splicing; protein product has enzymatic activity	Gerber et al. (1997)
Alu-J	Cathepsin B	Human	SRP RNA	Alt. splicing of exon 2 in 5' UTR	Berquin et al. (1997)
Alu	β 1C-2 integrin subunit	Human	SRP RNA	Alt. splicing, part of protein cod. region	Svineng et al. (1998)
Alu	DNA (cytosine-5) methyltransferase (CpG MTase)	Higher Primates	SRP RNA	Alt. splicing, part of protein cod. region	Hsu et al. (1999)
Alu	7.8 kb RNA	Human	SRP RNA	Induction of expression of a ST receptor in <i>trans</i>	Almenoff et al. (1994)
Alu	Wilms' tumor gene (WT1)	Human	SRP RNA	Intronic transcriptional silencer	Hewitt et al. (1995)
Alu	BRCA-1 gene, ERF-3	Human	SRP RNA	Estrogen-dependent transcriptional enhancers	Norris et al. (1995)
Alu	Parathyroid hormone gene	Human	SRP RNA	Negative calcium response element	McHaffie and Ralston (1995)
Alu	Poly(ADP-ribosyl) transferase (ADPRT) gene	Human	SRP RNA	Transcription regulation	Schweiger et al. (1995); Oei et al. (1997)
Alu	Potentially many genes	Human	SRP RNA	Transcriptional modulation via binding of YY1 protein	Humphrey et al. (1996)
Alu	Myeloperoxidase gene promoter	Human	SRP RNA	Composite SPI-thyroid hormone-retinoic acid response element	Piedrafita et al. (1996)
Alu	α 3 nicotinic receptor subunit	Human	SRP RNA	Transcription modulation	Fornasari et al. (1997)
RRE ^a	Erythropoietin recept. prom.	Mouse	?	Transcription inhibitor	Youssoufian and Lodish (1993)
Highly repet. element ^a	c-Ha-ras	Human	?	Blocks transcriptional readthrough	Lowndes et al. (1990)
MIR	Nicotinic acetylcholine receptor α subunit	Human	tRNA	Generating new splice variant, contributes to protein coding region	Murnane and Morales (1995)
MIR	β -tubulin	Human	tRNA	Polyadenylation signal	Murnane and Morales (1995)

Table 2. Continued

Retronuon	Gene/nuon that is under its influence	Species	Ancestor of source nuon	Serves as	References
MIR	Follitropin receptor	Sheep	tRNA	Polyadenylation signal	Murnane and Morales (1995)
MIR	Clone c-zrog02	Human	tRNA	Polyadenylation signal	Murnane and Morales (1995)
MIR	Clone NIB1273	Human	tRNA	Polyadenylation signal	Murnane and Morales (1995)
γ -actin	Salivary amylase gene	Human	mRNA	Promoter	Samuelson et al. (1990; 1996) Emi et al. (1988)

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Not all examples are proven exaptations. Especially events that date back not much more than a few million years could only be potential exaptations (potaptations according to Brosius and Gould, 1992, 1993).

^aResemblance to a known repetitive element not yet established.

Alu as selfish DNAs and does not require them ever to provide any useful function for the organism. An alternative model is the single source or 'master gene' model, envisaging Alu sequences as pseudogenes created by retroposition from a single functioning gene located at a specific locus This model hypothesizes that the source (or 'master') Alu is conserved by selection arising from its as-yet-unknown function." Much of the evidence favors the latter, master gene model (Willard, Nguyen & Schmid, 1987; Deininger & Slagel, 1988; Britten et al., 1988; Quentin, 1988; Labuda & Striker, 1989; Jurka & Milosavljevic, 1991; Deininger et al., 1992). The relationship of the first known master gene for a SINE, BC1 RNA, to ID repetitive elements (Kim et al., 1994; Deininger, et al., 1996) further supports this model. The findings on the interrelationship of BC1 RNA, its gene and related ID repetitive elements is further illuminating in that it regurgitates a frequently overlooked fact: black and white scenarios are relatively rare in biology; reality often lies between extremes (albeit sometimes much closer to one than to the other). BC1 RNA, generated by retroposition of a tRNA^{Ala}, has been exapted into a function in rodent nerve cells (Brosius & Tiedge, 1995). Depending on the rodent species, BC1 RNA is also the master gene for a few hundred up to 10,000 ID repetitive elements generated by retroposition. In some rodent species there is a large number of additional ID elements ($\geq 100,000$ in rat) whose consensus sequences point to a few different master genes. In the rat, up to four additional master genes most likely belong to the few ID elements that are actively being transcribed. For a more detailed discussion see (Brosius, 1999b).

It is quite rare that retroposition yields a novel transcribed gene encoding a small RNA, such as BC1 RNA, BC200 RNA (see below) or transcribed SINEs,

such as the postulated additional ID element master gene(s). Much more frequent is recruitment or exaptation of a SINE as new regulatory element for a targeted gene. This includes enhancement or silencing of transcription, providing (alternative) splice sites, polyadenylation signals, modulation of mRNA stability and small protein coding domains (Table 2).

Why do many SINEs contain LINE sequences at their 3' end? Template switching as alternative explanation

SINEs and other retrosequences depend on LINEs and/or other retroelements to provide for the key enzyme reverse transcriptase ((Feng et al., 1996); see also below). In addition, LINEs have been suggested to play a further role in the propagation of SINEs (Ohshima et al., 1996; Okada et al., 1997) as well as mRNA derived retrosequences (Jurka, 1997). It is thought that a strong-stop DNA (generated by reverse transcription) with a primer tRNA was integrated into the 3' portion of a LINE, which gave rise to a primordial SINE. Here, I represent an alternative explanation with an equally tight association of a LINE to a tRNA – reverse transcription of a LINE is aborted (presumably again through a strong stop), and then the template switched (Pathak & Hu, 1997) to a nearby (primer) tRNA or other small RNA, possibly co-packaged in virus-like particles (Kolosha & Martin, 1995). Some of these SINEs with 5' tRNA domains and 3' LINE domains – if transcribed – could be efficient founder genes for additional SINEs, presumably by providing more efficient priming sites for reverse transcriptase. However, not all SINEs feature this composite structure, notable exceptions being Alu, B1, B2, ID and S1 repetitive elements. Hence, alternative features must exist that earmark small RNAs as efficient master genes.

Table 3. Vertebrate genes generated by retrosequences

Retrogene; Pattern of expression; chromosome	Species	Source gene; pattern of expression; (# of introns); chromosome	Template for reverse transcription	Hallmarks of retrosequences			References
				intron loss	A-stretch at 3' end of foun- der RNA	direct repeats	
Insulin I; Langer- hans islets; (1)	Murids	Insulin II; Langer- hans islets; (2)	Part. proc. hnRNA	(+)	+	(+)	Soares et al. (1985)
Phosphoglycerate kinase (Pgk-2); testes	Mammals	Pgk-1; constitu- tive; (10); X-linked	Mature mRNA	+	(+)	(+)	McCarrey and Thomas (1987); Boer et al. (1987); Adra et al. (1988)
Zfa; testes; chr 10	Mouse	Zfx; ubiquitous; (≥ 10); X-linked	Mature mRNA	+	(+)	+	Ashworth et al. (1990)
Pyruvate dehydro- genase (Pdha2); testes	Human mouse	Pdha1; constitutive; (10); X-linked	Mature mRNA	+	(+)	(+)	Dahl et al. (1990) Fitzgerald et al. (1992)
N-myc2; brain and liver tumours ^a	Sciuridae rodents, e.g. woodchucks	N-myc1; in development and various adult tissues; (2)	Mature mRNA	+	(+)	+	Fourel et al. (1990, 1992); Sugiyama et al. (1989, 1999); Robertson et al. (1991); Quigon et al. (1996)
NB-1 or CLP; epithelial tissue; chr 10	Human	Calmodulin CaMIII; ubiquitous; (5)	Mature mRNA	+		(+)	Yaswen et al. (1992); Rhyner et al. (1992); Berchtold et al. (1992)
Carcinoma associated anti- gen, GA733-1	Human	GA733-2; placenta, carcinoma; (8)	mRNA	+			Linnenbach et al. (1989, 1993)
Glutamate dehydrogenase (GLUD2); retina, testes, brain; X-linked	Human	GLUD1; ubiquitous; (13); chr 10	Mature mRNA	+	+	+	Shashidharan et al. (1994)
S-adenosyl- methio- nine decarboxylase (AMD2); liver and other tiss.; chr 12	Mouse	(AMD1); ubiquitous; (9)	Mature mRNA	+	+	+	Persson et al. (1995, 1999); Nishimura et al. (1998)
Glucose-6-phosphate dehydrogenase (G6PD-2); testes	Mouse	G6PD-1; constitutive (10); X-linked	Mature mRNA	+	+	+	Hendriksen et al. (1997)
Hypoxanthine transferase, <i>HPRT</i> -2; liver	Kangaroo	<i>HPRT</i> -1; ubiquitous; (8); X chr	Mature mRNA	+		(+)	Noyce et al. (1997); Noyce and Piper (1994)

Table 3. Continued

Poly(A) binding protein 2 (Pabp2); spermatogenic cells	Mouse	Pabp1; spermatogenic and somatic cells (several)	Mature mRNA	+			Kleene et al. (1998)
C γ catalytic subunit of cAMP-dependent protein kinase ^b ; testes; chr 9	Catarrhini primates	C α catalytic subunit of cAMP-dependent protein kinase; ubiquitous; (~9); chrom. 19	Mature mRNA	+	(+)	(+)	Reinton et al. (1998)
H430 encoding a splicing factor; pancreas, spleen, prostate etc.; chr 11	Human	PR264/SC35; thymus, spleen, kidney, lung etc.; (2); chr 17	Mature mRNA	+	(+)	+	Soret et al. (1998)
CDY genes (at least one family member); testes; Y chr	Anthropoidea	CDYL; ubiquitous; (9); chr 13 (human)	Mature mRNA	+			Lahn and Page (1999)
Proteasome activator PA28, β -subunit (PMSE2b); constitutive; chr 14	Mouse	PMSE2, gamma interferon inducible; (10); chr 11	Mature mRNA	+	+	+	Zaiß and Kloetzel (1999)
Centrin, <i>Cetm1</i> ; testes; chr 18	Mammals	<i>Cetm2</i> ; neonatal testes, oviduct; (4); X chr	Mature mRNA	+		+	Hart et al. (1999)
XAP-5-like (X5L) ^c ; chr 6	Human	XAP-5; (12); X-linked	Mature mRNA				Sedlacek et al. (1999)
BC1 RNA; neurons; chr 7 (mouse)	Mouse						
	Rodents	tRNA ^{Ala} ; ubiquitous	Non-mRNA	n.a.	+	(+)	DeChiara and Brosius (1987); Martignetti and Brosius (1993a)
BC200 RNA; neurons; chr 2 (human)	Anthropoidea	Free Alu monomer	Non-mRNA	n.a.	+	(+)	Martignetti and Brosius (1993b)

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^aSee also s-myc, sm-myc in rodents and mycL2 in primates; role in apoptosis.

^bProtein product not confirmed yet.

^cIntron in 5' UT.

The role of SINEs in the evolutionary roulette

It has been argued before that the shotgunning of mobile regulatory elements through the genome is a powerful way to impose plasticity to the genome (McClintock, 1948; Georgiev, 1984; Brosius & Gould, 1992). Kermekchiev et al. (1991) have shown that various combinations of *in vitro* shuffled promoters and enhancers function with high efficiency. The authors proposed, therefore, that "a generally permissive enhancer/promoter interaction is of evolutionary benefit for higher eukaryotes: by enhancer shuffling, genes could be easily brought under a new type of inducibility/cell-type specificity". Retronuons as shown in Tables

1 and 2 are precisely the mobile elements that could have achieved and still accomplish a constant shuffling of regulatory elements on a trial-and-error basis.

The essence of the human/chimp divergence at the genomic level

A comparison between genomes of closely related species will reveal the driving forces of speciation. Especially the full sequences of human and chimpanzee will disclose that there are few genes that arose *de novo* in one of the two species. Naturally, we expect some genes in one species versus the other to have arisen by duplication (via recombination or

Table 4. Vertebrate genes probably generated by retrosequences

Retrogene; pattern of expression; chromosome	Species	Presumable source gene; pattern of expression; (# of introns); chromosome	Template f. reverse transcript.	Age of retrogene	References
Replication-dependent histone genes ^a	Various	'Replacement' variant histone genes	mRNA	Metazoans and plants	Reviewed in: Kedes et al. (1979); Hentschel and Birnstiel (1981) Wilkie et al. (1992)
G protein α subunit, Gi class, Gnaz	Mammals	Gnai; (8)	Part. proc. hnRNA		
G-protein coupled receptors ^b	Various	G-protein coupled receptors	mRNA		Reviewed in: Gentles and Karlín (1999); Brosius (1999)
Potassium channels	Various	Potassium channel	mRNA		Reviewed in: Strong et al. (1993)
Class III POU domain proteins e.g. SCIP (or Tst-1, Oct-6); or: Brn-3b; brain; X chr; early development and brain	Various	POU domain transcription factor; (multiple) Brn-3a or 3c; brain; chr 14 and chr 18	mRNA		Kuhn et al. (1991); Hara et al. (1992); Theil et al. (1994); Alvarez-Bolado et al. (1995); Atanasoski et al. (1995); Levavasseur et al. (1998)
Forkhead transcription factors, e.g.: brain factor-2 (HBF2) ^c ; fetal brain; chr 14; MFH-1	Various	Brain factor-1 (HBF1); fetal brain; (1) chr 14	mRNA		Wiese et al. (1995); Ernstsson et al. (1996); Frank and Zoll (1998); Miura et al. (1997)
Inducible heat shock genes ^d	Various	Constitutive heat shock genes	mRNA	Ancient	Hunt and Morimoto (1985); Mues et al. (1986); Zakeri et al. (1988); Milner and Campbell (1990); Lim and Brenner (1999)
Genes encoded by herpesviruses	Various	Var. cellular intron- containing genes	mRNA		Reviewed by Brunovskis and Kung (1996); Martin (1999)
Protamines	Vertebrates	Protamines			States et al. (1992); Jankowski et al. (1986); Moir and Dixon (1988); Oliva and Dixon (1989); Retief et al. (1993); Schlüter and Engel (1995)
Non-histone chromosomal protein HMG-1	Mammals	HMG-2; (4)	mRNA	\geq mamm. radiation	Stros and Dixon (1993); Stros et al. (1995)
Glycerol kinase (GyK) ^e ; testes; chr 4	Human	GyK; constitutive; (18);	mRNA		Sargent et al. (1994b)
Antioxidant protein 2 related seq. (Aop2-rs1 and Aop2-rs2 ^f); var. tissues; chr 2 and chr 4, respectively	Mouse	X-linked Antioxidant protein 2 (Aop2); var. include. heart, liver, kidney; (4); chr 1	mRNA		Pan et al. (1999) Phelan et al. (1998)
DNA ligase IV (LIG4)	Mouse		mRNA		Barnes et al. (1998)
Actin-like-7A and actin-like-7B (<i>ACTL7A</i> , <i>ACTL7B</i>); chr. 9	Mammals	Actin or an actin-related protein (ARP)	mRNA		Chadwick et al. (1999)
Sterol 12 α -hydroxylase (<i>CYP8B1</i>); 3p21.3 (Hsa), 9qF4 (Mmu)	Human Mouse	<i>CYP8A1</i> (or <i>CYP7A1/7B1</i>)	mRNA	\geq mamm. radiation	Gäfvéls et al. (1999)

Table 4. Continued

Metalloproteinase-distintegrins (ADAM20, ADAM21)	Human	Intron-containing family members	mRNA	Poindexter et al. (1999)
α CP-1 RNA binding protein	Mammals	α CP-2; (≥ 12)	mRNA	Makeyev et al. (1999)
Germ cell-specific actin capping protein α (Gsg3 clone); chr 6	Rodents	Somatic cell type actin capping protein α (ACP α)	mRNA	Yoshimura et al. (1999)
1-Cys peroxiredoxin, 1-Cys Prx (CP-2 and CP-5)	Mouse	CP-3; (4)		Lee et al. (1999)

Intron loss in comparison to an introncontaining paralogue candidate is the only (remaining) hallmark. The decision, whether a sequence belongs in this table or Table 3 is arbitrary in some cases. Likewise, inclusion in Table 4 versus Tables 5 or 6 is somewhat arbitrary. In situations where only one intron is present in the putative founder gene, it may have been acquired in the founder. (e.g. protamine genes). Clearly, not all examples can be proven exaptations (potaptations according to Brosius & Gould, 1992, 1993).

^aWe cannot rule out that the ancestral histone gene was intronless and some histone genes acquired introns in the 'intron late' scenario.

^bMany genes encoding G-protein coupled receptors that lack introns in their coding regions feature an intron in the 5' UT; presumably generated by acquisition of splice sites (Brosius & Gould, 1992).

^cIntronless HBF-2 is clustered with HBF-1 (one intron in coding region) on chromosome 14q 11-13 (Wiese et al., 1995).

^dA 71 kDa heat shock protein has been described in the human genome that contains 8 introns (Dworniczak & Mirault, 1987).

^eIn addition to the split gene there are at least six additional loci in humans; two are pseudogenes (Xq and chr. 1) two are active retrogenes (both chr. 4) - protein product not confirmed yet; the status of the remaining two genes needs to be established.

^fAop2-rs2 potentially encodes only a truncated polypeptide of 114 aa.

retroposition) after the split. Even then, five million years do not provide enough time to generate much sequence divergence. I predict, that the essential difference between these closely related primate species is not the emergence of a certain number of novel genes. Instead, we will find that the main variance will be observed:

- (i) in the recruitment of short domains of protein sequences from anonymous sequences (or retronuon-derived sequences) previously located in introns (Gilbert, 1978) or intergenic sequences and
- (ii) most importantly, in the differential expression of shared genes.

These changes of when and where genes are being expressed with numerous consequences (including governing different developmental processes) will be found to be largely determined by migrant retronuons.

Tip of the iceberg as many regulatory elements could have been derived from ancient retronuons

Clearly, Tables 1 and 2 exhibit only the tip of the iceberg for a number of reasons – once the entire sequences of genomes of higher Eucarya will become available, many more examples will become evident. Although juxtaposition of retronuons with known and unknown genes will become apparent by sequence analysis, the question remains whether such retronuons do have an effect on the targeted gene. A

global approach in the postgenomic era is needed to identify most retroelements that exert an effect on associated genes. There is yet another important reason why even then these tables would remain incomplete to a large degree: on an evolutionary scale, there is only a relatively small time window when retronuons as shown in Tables 1 and 2 are discernible. In other words, about 200×10^6 years from now the retronuon from Table 1 or 2 that carried an enhancer to an existing gene will have disintegrated. What remains, for example, are short conserved enhancer sequence motifs, perhaps after some modification of the enhancer core itself. It is clear that retroposition not only took place in the last 200×10^6 years but probably since the transition from the RNP to the DNA worlds (Darnell & Doolittle, 1986; Flavell, 1995; Brosius, 1999c; Poole Jeffares & Penny, 1999). Therefore, it would be of little surprise if it turned out that the vast majority of such small regulatory motifs that today are visible and active near targeted genes were derived from ancient retronuons whose identities disappeared over time (Brosius & Gould, 1992); W. Herr, cited in (Brosius & Tiedge, 1996), (Britten, 1996, 1997).

Messenger RNA derived retrosequences jump too. Did all intronless genes arise via an RNA intermediate?

Just as regulatory elements can insert next to resident genes, retroposition also allows for the opposite

Table 5. Intronless* vertebrate genes (no further evidence of retrosequence origin)

Gene	Species	Pattern of expression	References
Interferons	Vertebrates		Reviewed in: Nagata et al. (1980); Lawn et al. (1981); Watkins et al. (1991); Roberts et al. (1998)
Ribonucleases ^a	Various		Carsana et al. (1988); Hamann et al. (1990); Samuelson et al. (1991); Sasso et al. (1991); Tiffany et al. (1996)
Mos	Mammals		Watson et al. (1982); Newman and Dai (1996)
Insulinoma associated, IA-1 (zinc finger); chr 20	Human	Neuroendocrine tumours	Lan et al. (1994); Li et al. (1997)
Transcription elongation factor SII or A (TCEA)	Human		Park et al. (1994); DiMarco et al. (1996)
Modifier of Na ⁺ -D-glucose co-transport (hRS1)	Human		Lambotte et al. (1996)
Profilaggrin	Mouse, rat		Haydock and Dale (1986)
Thrombomodulin	Mammals	RA inducible	Jackman et al. (1987); Niforas et al. (1993)
HS, HGT-C2, HGT-B2, BIIIB4, HGT-F, cKer1 keratins; keratin-associated protein, Krtap12-1; KAP6; B2E and B2F (high sulfur protein genes, in hair follicles); keratin-associated proteins pmg-1 and pmg-2	Vertebrates	Hair, skin	Powell and Rogers (1986); Kuczek and Rogers (1987); Frenkel et al. (1989); Whitbread et al. (1991); Fratini et al. (1993); Mitsui et al. (1998); Cole and Reeves (1998); Kuhn et al. (1999)
Blood platelet membrane glycoprotein Ib α , glycoprotein V (GPV); glycoproteins Ib β , IX ^b	Mammals	Platelets	Wenger et al. (1988); Lanza et al. (1993); Ravanat et al. (1997); Yagi et al. (1995)
Olfactory marker protein (OMP)	Rat		Danciger et al. (1989)
Melanin-concentrating hormone	Fish		Takayama et al. (1989)
Cerebellar degeneration-related antigen, CDR34	Human		Chen et al. (1990)
Leukosialin CD43	Mammals		Cyster et al. (1990); Shelley et al. (1990)
Nuclear pore glycoprotein p62	Rat		D'Onofrio et al. (1991)
N-acetyltransferases Nat1 and Nat2 ^a	Vertebrates		Grant et al. (1989); Blum et al. (1990a-c); Martell et al. (1991)
Centromere protein, CENP-B	Mammals		Sullivan and Glass (1991); Bejarano and Valdivia (1996)
JUN protooncogene	Vertebrates		Hattori et al. (1988); Hartl et al. (1991)
Factor VIII-associated gene (F8A) ^d	Mammals	Ubiquitous	Levinson et al. (1992)
A-kinase anchor protein, AKAP 75	Bovine		Hirsch et al. (1992)
LAP, C/EBP α , β ₁ , δ ₁ , CRP2 or NF-IL6 β	Human		Landschulz et al. (1988); Akira et al. (1990); Chang et al. (1990); Descombes et al. (1990); Cao et al. (1991); Williams et al. (1991); Kinoshita et al. (1992)
CAAT/enhancer-binding proteins; basic region-leucine zipper class (bZIP)			
Cytochrome b5	Rabbit		Takematsu et al. (1992)
Gap junction genes connexin 31.1 and 30.3; chr 4	Mouse	Skin	Hennemann et al. (1992)
Na ⁺ -MI cotransporter (SMIT/SLC5A3)	Human	Kidney and other tissues	Berry et al. (1995); Porcellati et al. (1999)
Myeloid zinc finger gene (MZF-1)	Human	Bone marrow	Hui et al. (1995)
U2 auxiliary factor binding protein related sequence	Human		Pearsall et al. (1996)
U2AFBPL ^c chr 5 / U2afbp-rs	Mouse		

Table 5. Continued

(imprinted in mouse); chr 11			
Acetyltransferases AT1 and AT2	Rat	Various	Land et al. (1996)
Choriolysin H (HCE)	Teleost fish		Yasumasu et al. (1996)
Pw1 zinc-finger protein			Relaix et al. (1996)
Defensin (HNP-1)	Human		Takemura et al. (1996)
Rho/Rac-like RhoG GTPase ^f (ARHG)	Human		Le Gallic and Fort (1997)
Ventral prostate protein C7orf1	Human		Peacock et al. (1997)
Antiproliferative proteins Tob, ANA	Mammals	Ubiquitous	Yoshida et al. (1997, 1998); Guéhenneux et al. (1987)
Serine-threonine kinase genes Tsk1, Tsk2	Mouse		Galili et al. (1997)
Glycosylphosphatidylinositol synthesis gene PIGC			Hong et al. (1997)
Growth arrest-specific C16orf3	Human		Whitmore et al. (1998)
Sex determining gene SRY and SOX-3	Mammals		Foster and Graves (1994); Tucker and Lundrigan (1995); O'Neill et al. (1998)
2', 5'-oligoadenylate-dependent RNase (interferon inducible)	Human		Tnani and Bayard (1998)
Necdin	Mammals	Neurons	Uetsuki et al. (1996); Nakada et al. (1998)
Chondroitin 6-sulfotransferase (C6ST); chr 11	Human		Mazany et al. (1998)
Testes-specific protein	Mammals	Ubiquitous	Vogel et al. (1998)
Y-encoded-like, (human TSPYL chr 6; rodent Tspyl chr 10)			
Citrate synthase (CS); chr 12	Human		Goldenthal et al. (1998)
CXorf1	Human	Hippocampus	Redolfi et al. (1998)
Cholesterol 25-hydroxylase	Human, mouse		Lund et al. (1998)
Prion protein ^g	Mammals, chicken		Lee et al. (1998)
Insulin receptor substrate 4 (IRS-4) ^h	Mouse		Fantin et al. (1999)
110 kDa high molecular wt. basic nuclear protein (HMrBNP)	Flounder	Sperm	Watson and Davies (1999)
cded/lor	Mouse		Mishra et al. (1999)
Rab-like protein (Rlp-2)	Human		Peng et al. (1999)
Slow-kinetics immediate early gene ler5	Mouse		Williams et al. (1999)
transport modifier RS1	Rabbit		Reinhardt et al. (1999)
Sperizin, RING zinc-finger protein	Mouse	Haploid sperm cells	Fujii et al. (1999)
Malaria-inducible gene	Mouse	Spleen	Krücken et al. (1999)
KRML (MAFB); chr 20	Human	Hemopoietic tissue	Wang et al. (1999)
α -endosulfine (ENSA); chr 14	Human		Heron et al. (1999)
ZNF127 RING zinc-finger protein ⁱ	Human		Jong et al. (1999)
MAGEL2 ⁱ	Human, mouse	Brain, placenta	Boccaccio et al. (1999)

*May have intron(s) upstream from coding region; a single intron in the coding region may also have been generated subsequent to retroposition (e.g. the monocyte-specific Dif-2 gene (Pietzsch et al., 1998) or the acidic 80 kDa protein kinase C substrate/MARCKS (Erusalimsky et al., 1991; Blackshear et al., 1992).

^aSeveral genes contain single introns in 5' UT. ^bNo intron in entire ORF, but 5' UT. ^cCoding region contains a tandem hexapeptide repetitive structure; could have been exapted from a non-coding repeat region. ^dOne of the human genes may be located in intron of factor VIII gene.

^eOne of the human genes located on the X chromosome contains introns (Kitagawa et al., 1995). ^fContains a large exon in the 5' UT. ^gMouse, sheep have two and humans one exon(s) in 5' UT. ^hRelated gene IRS-3 contains one intron in the coding region. IRS-1 and IRS-2 are thought to be intronless as well. However, Vassen et al. (1999) described an intron at the C-terminus of the IRS-2 ORF. ⁱLocated on chr 15q11-13 (Prader-Willi Syndrome, PWS, region); paternally expressed.

scenario: retrogenes, usually generated by reverse transcription of mRNAs, can insert next to resident promoter/enhancer elements (Brosius, 1991) and thus escape transcriptional silencing (the fate of most retrosequences). The retrogene is often expressed in a cell type and/or at a stage different from the founder gene. In many cases a retrogene, when different from its founder gene by modifications, can be recruited or exapted into a novel function (Table 3). In fact, probably the majority of intronless genes may have arisen by this mechanism although most of their hallmarks may not be recognizable anymore (Tables 4, 5). This would be certainly true if the 'intron-early' hypothesis was correct. Should the 'intron-late' camp be correct, at least some of the intronless genes may be ancient and have remained intronless from 1.7 to 1.0×10^9 years ago, the presumed period when spliceosomal introns arose (Cavalier-Smith, 1991), until now. Apart from the scenario that a retrogene is immediately active upon integration, there are scenarios where the retrogene is initially inactive and is being activated at a later time, for example, upon activation of flanking regulatory elements by mutation or acquisition of a retronuon.

Inactive retrogenes: future genes in the making?

It is conceivable that many retrogenes were reactivated after a considerable amount of transcriptional and/or translational latency. This time of inactivity facilitated more drastic changes in gene structure without the possible 'danger' of expressing intermediate molecules that had a large degree of similarity to the original ones but exerted detrimental effects as is the case in dominant negative mutations. As early as 1972, Arthur Koch (1972) had recognized the importance of untranslatable intermediates in enzyme evolution (for a more detailed discussion, see Brosius, 1999a, b).

Virtually any retrogene listed in Table 6 may be in such a 'stand-by' mode. One may say that even the majority of retrogenes, so-called inactive retroseudogenes, their ORFs annihilated with indels and mutations that introduce many stop codons in the coding region may be 'on call' (potaptation = potential exaptation) for future utility (exaptation) (Gould & Vrba, 1982). In summary, the relentless activity of retronuons in most vertebrate genomes had and will continue to have important consequences for the evolution of populations and species (Rose & Doolittle, 1983, 1989; McDonald, 1990, 1995).

Keeping retroposition in check

Countermeasures of the host

Despite potential advantages of retroposition on future evolvability, if a host 'permits' unrestricted retroposition, 'runaway' activity may result. Left without restraint, this may lead to disadvantages such as frequent inactivation or misregulation of essential genes and overinflation of genome size. It has been suggested, therefore, that the host is devising countermeasures to keep retroposition in check. For example, Patience wrote: '... animals harboring infectious ERVs have evolved mechanisms to prevent re-infection with their own potentially infectious ERV genomes' (Patience et al., 1997). This is perhaps an analogy to the arms race between host defense and external parasites (microorganisms/viruses). This analogy also includes gradual reduction of the parasite's virulence in order to ensure its own survival. It has been proposed that a widespread defensive measure of the host against retronuons is methylation of retroelements in order to shut down their transcription (Walsh et al., 1998; Wolffe & Matzke, 1999). While such measures may be effective against retroelements such as proviruses or intact LINEs, the methylation of truncated LINEs or SINEs such as Alu elements seems futile, since, for example, very few of the 10^6 Alu repeats are transcriptionally and hence retropositionally active. Likewise, promotion of mutations at methylated CpG sites is of little consequence for the propagation of Alu elements. In the case of SINEs, the only effective countermeasure would be silencing of the few active master genes. This, in turn, may be impossible if the RNA product of the master gene is under selective pressure. As a more effective measure, the activity of reverse transcription could be blocked provided there are no negative consequences for the host (see below). As a result, production of retroelements and retrosequences alike would be stopped, the only way 'out' for a parasitic retroelement being a change of host by horizontal transfer (Jordan et al., 1999).

Parasites or symbionts

Despite the realization that occasionally a TE is advantageous to the host, many authors consider transposable elements (TEs) including retronuons as genomic parasites (Doolittle & Sapienza, 1980; Orgel & Crick, 1980; Hickey, 1982; Kidwell & Lisch, 1997; Walsh, Chaillet & Bestor, 1998; Jordan, Matyunina

Table 6. Intronless vertebrate genes likely of retroposition origin – no proven activity of gene product (this does not exclude transcriptional or even translational activity)

Retrogene; pattern of expression; chromosome	Species	Source gene; pattern of expression; (# of introns); chromosome	Age of retrogene	References
Non-muscle tropomyosin (hTM _{NM} -1)	Human			MacLeod et al. (1983)
Metallothionein (MT-1 Ψ b) ^a	Rat			Andersen et al. (1986)
Sarcomeric actin α 2	Frog	Actin		Stutz and Spohr (1987)
Glutamine synthetase (GSr)	Mouse	Glutamine synthetase (GSi)		Bhandari et al. (1991)
Glutamine synthetase (Ψ GS) ^b	Human	Glutamine synthetase		Chakrabarti et al. (1995)
Heat stable antigen (2 ORFs)	Mouse	Heat stable antigen		Wenger et al. (1991)
Adenylate kinase 3 (AK3) ^c ; chr 17	Human	AK3; chr 9		Xu et al. (1992)
Ferritin L subunit Lg	Mouse	ferritin L subunit; (3)		Renaudie et al. (1992)
Processed CD-MPR gene ^d ; chr 3	Mouse	Cation-dependent mannose 6-phosphate receptor (CD-MPR); (7); chr 6		Ludwig et al. (1992)
Id2B ^e	Human	Helix-loop-helix protein Id2		Kurabayashi et al. (1993)
Casein kinase II α	Human	Casein kinase II α		Devilat and Carvallo (1993)
Ψ EF1 _A #1 ^f	Bovine	CCAAT transcription factor subunit EF1 _A		Ozer et al. (1993)
Ψ 5HT1D ^g	Human	serotonin receptor 5HT1D		Bard et a. (1995)
Protein kinase C (Ψ PKC ζ ^h)	Rat	PKC ζ		Andrea and Walsh (1995)
FAU1P ⁱ ; chr 18	Human	FAU1		Kas et al. (1995)
dbpB pseudogene ^j	Human	DNA binding protein dbpB		Kudo et al. (1995)
mif rp-1	Mouse	macrophage migration inhibitory factor (MIF)		Bozza et al. (1995)
Ferritin H subunit pseudogene	Human	Ferritin H subunit		Zheng et al. (1995, 1997)
Prothymosin α intronless	Mammals	prothymosin α		Varghese and Kronenberg (1991); Manrow et al. (1992); Rubtsov and Vartapetyan (1995)
Laminin receptor (37LRP/p40), intronless	Human	laminin receptor (37LRP/p40)		Jackers et al. (1996)
LAMRL5 ^k	Human	67-kDA laminin receptor (LAMR1)		Richardson et al. (1998)
Ubiquitin-conjugating enzyme UBE2L1; chr 14	Human	ubiquitin-conjugating enzyme UBE2L3; (1); chr 22		Moynihan et al. (1996)
Ψ Adh-2 ^l	Mouse	Class III alcohol dehydrogenase (Adh-2); (8)		Foglio and Duester (1996)
MSSP-1 (transcriptional enhancer of c-myc)	Human	MSSP-2; (15)		Haigermoser et al. (1996)
Olfactory receptor pseudogenes	Human	Olfactory receptor genes		Crowe et al. (1996)
Hp53int1 ^m	Human			Reisman et al. (1996)
Phosphoglycerate mutase brain isoform	Human	Phosphoglycerate mutase brain isoform (<i>PGAM1</i>)		Dierick et al. (1997)
pseudogene (Ψ PGAM1 ⁿ)				
α tubulin-related sequence; chr 11 ^o	Human	Keratinocyte α tubulin		Devon et al. (1997)
r.pem2 homeobox gene ^p ; epipydimis; X- linked	Rat	r.Pem, orphan homeobox gene; testes, ovary, placenta, epididymis; (5); chr 4		Nhim et al. (1997)
Leukocyte antigen C1pg-26 ^q	Dog	Leukocyte antigen DLA class I; (7)		Burnett et al. (1997)

Table 6. Continued

Tdgl1-ps1; chr 16	Mouse	Teratocarcinoma-derived growth factor- 1; Tdgl1; (5); chr 9	Liguori et al. (1996, 1997)
TDGF3; X chr	Human	TDGF1; chr 3	Dono et al. (1991)
(FABP3-ps); chr 13	Human	Fatty acid binding protein FABP3, chr 1	Prinsen et al. (1997)
Serotonin-7 receptor (5-HT7 Ψ) ^f	Human	5-HT7	Quian et al. (1998)
Mannose-binding protein-A; chr 10	Human		Guo et al. (1998)
Ψ FGFR-3 (partial, antisense); fetal development; chr 1	Mouse	Fibroblast growth factor receptor (FGFR-3); chr 5	Weil et al. (1997)
Ψ ribosomal protein L7 (antisense)	Human	Ribosomal prot. L7	Hohlbaum et al. (1998)
Supt4h2; chr 10	Mouse	Supt4h; (4); chr 11	Chiang et al. (1998)
Ubiquitin conjugating E2 enzyme ubc9-ps1 and ubc-9-ps12	Mouse	ubc-9	Tsytsykova et al. (1998)
SMT3A and 3 SMT3B proc. pseudogenes	Mouse	Ubiquitin-like proteins	Chen et al. (1998)
Ψ PTEN ^s , chr 9	Human	PTEN/MMAC1/TEP1 phosphatase; chr 10	Dahia et al. (1998)
EIF4E2 translational initiation factor	Human	EIF4E1; (6)	Recent Gao et al. (1998)
<i>EIF2γ</i> A; testes; chr 12	Human	Euk. Translation initiation factor eIF-2 γ (<i>EIF2γ</i> X); X-linked	Ehrmann et al. (1998)
Ψ hGABP α t	Human	ets related GAPB α	Luo et al. (1999)
Proto-oncogene hPTTG2	Human	hPTTG1	Prezant et al. (1999)
CDC42-like; chr 4	Human	CDC42; chr 1	Nicole et al. (1999)
Spondyloepiphyseal dysplasia tarda gene (SEDLP)5; many tissues; chr 19	Human	Spondyloepiphyseal dysplasia tarda gene (SEDLP or GPM6B); many tissues; (3); X-linked	Recent Gedeon et al. (1999)
CK2 α ; chr 11	Human	CK2 α ; (12); chr 11	Wirkner and Pyerin (1999)

There are probably numerous additional retrogenes whose ORFs are not severely compromised or could yield a truncated polypeptide, partially in a different reading frame (e.g. Chen et al., 1982; Varshney & Gedamu, 1984; Dudov & Perry, 1984; Nojima et al., 1987; Kuzumaki et al., 1987; Srikantha et al., 1987; Seelan & Padmanaban, 1988; Nielsen & Trachsel, 1988; Kawaichi et al., 1992; Jun et al., 1997; Palmer et al., 1998). However, transcription and/or translation are not documented.

^aThis retrosequence is transcribed. Due to an insertion after codon 28 the ORF is shifted to with a different hypothetical C-terminus of an additional 35 amino acids instead of 33 aa in the correct MT-1 frame.

^bThis retrosequence is transcribed. The ORF is truncated but retains $\sim 2/3$ of the coding sequence; probably no protein product.

^cEmbedded in intron 10 of NF1 gene (located on human chr 17).

^dThis retrosequence is transcribed. The ORF is truncated after 141 codons (out of 278 possible) in murine CD-MPR. A soluble truncated form of CD-MPR encoding only the *n*-terminal extracytoplasmic region including codon 154 was functional in ligand binding and acid-dependent dissociation (Marron-Terada et al., 1998).

^eThis retrosequence is transcribed. Stop codon at aa 37, however.

^fThis retrosequence is transcribed. The ORF is truncated; probably no protein product.

^gThis retrosequence hypothetically encodes a 140 aa polypeptide most of which (aa 31–140) are similar to bovine EFl_A (324 aa total).

^hThis retrosequence is transcribed specifically in the brain. The ORF is truncated and no protein product could be identified by Western blots.

ⁱThis retrosequence is not transcribed but contains an intact ORF.

^jOne of 16 pseudogenes contains an intact ORF.

^kPotentially active retrogenes may also exist in the mouse (Bignon et al., 1991).

^l25 point mutations relative to Adh-2 cDNA, nevertheless ORF is intact, but no evidence for transcription, thus far.

^mLocated in the 10 kb first intron of p53 tumour suppressor gene; no or short ORF.

ⁿLocated in intron 1 of Menckes disease gene (*ATP7A*, *MNK*).

^oThe ORF is truncated but retains 80% of the coding sequence.

^pAlthough a processed mRNA was the founder of this retrogene, it acquired new splice sites that remove three premature stop codons yielding again an open reading frame – protein product not confirmed yet.

^qContains single ORF of 332 codons but no potential start codon in the *N*-terminal 2/3 of ORF; not likely to be functional.

^rTranscribed but not translatable.

^sORF intact, hypothetical polypeptide somewhat smaller due to loss of first start codon; no evidence for transcription as of yet.

^tThis retrogene is transcribed in human myeloid cells, but a mutation at the site that corresponds to the ATG start methionine codon may prevent its translation.

Table 7. Intron containing vertebrate genes featuring large exons (probably of retrosequence origin)

Gene	Species	Pattern of expression	References
Developmentally regulated type X collagen	Chicken		Ninomiya et al. (1986)
<i>Clr</i> and <i>Cl</i> s complement	Human		Tosi et al. (1989)
Follicle-stimulating hormone receptor (FSHR); LH, TSH receptors	Human		Gromoll et al. (1996); Misrahi et al. (1996)
Islet homeobox gene (<i>isl 1</i>) ^a	Mammals		Bozzi et al. (1996)

^aFeatures intronless homeobox domain.

& McDonald, 1999). Personally, I prefer the term genomic symbionts for a number of reasons: The translation of 'symbiosis' from Greek (living together) is at least as close to TE reality as is 'parasitiasis' (living at one's cost). G.P. Georgiev (1998) who was among the first to realize the biological significance of mobile genomic elements considered them not as 'pure genomic parasites' but as 'genomic symbionts.' He continues, 'Their importance is in spreading signal sequences of different type throughout the genome to supply different genetic elements with such signal sequences.... One may speculate that mobile elements are, at least sometimes, involved in regulation of certain gene expression, RNA splicing, replication initiation etc.... the existence of mobile elements sharply increases the variability and, in this way, accelerates the process of evolution. Mobile elements can make organisms fit more readily to changing environmental conditions. Therefore, the advantages gained from their presence may outweigh the expenses spent for their replication and expression' (Georgiev, 1984). It is remarkable that Georgiev had this insight one and a half decades prior to gradual acceptance of such concepts within the scientific community.

As mentioned earlier, the truth hovers most of the time in between extremes. Who would place smallpox, measles, influenza, typhus, bubonic plague and other infectious disease causing microbes not into the category of human parasites? Yet, these infectious agents that had 'plagued' European populations for centuries (implementation of host-infective agent strategies reduced the virulence of some over time) gave initially a handful of 'conquistadores' a selective advantage over the natives of the Americas by killing as much as 95% of the population (Diamond, 1997). Therefore, it is not so straightforward anymore whether germs belong to the parasitic corner, when the evolution of populations is considered. Likewise, it is conceivable that host interactions have blunted the negative impacts of TEs including retronuons (Kidwell & Lisch,

1997). The impact of retronuons on individuals, populations or species is to be considered separately (Doolittle, 1989). What may be a parasite for an individual, may be a symbiont for a population or species.

Genome expansion of retronuons lessens their negative impacts

Apart from the aforementioned reduction of transpositional activity by methylation and other measures (Kidwell & Lisch, 1997) genome expansion is an effective way to reduce negative impacts of retronuon insertion. By blowing up the size of introns and intergenic regions, the chances that a retronuon hits a 'vulnerable' target (exon, promoter, or other regulatory element) is greatly reduced. This means that yeast and *Drosophila*, for example, are more assailable to TE activities with negative consequences as are plants (such as maize) or mammals. It will be interesting, for example, to compare retroposition activity in species with tightly packed genomes such as the pufferfish (*Fugu rubripes*) with similar, more bloated, sequences. The *Fugu* genome comprising ~400 Megabases is about four-fold compressed in comparison to fish with larger introns and intergenic regions (zebrafish, *Danio rerio*, ~1700 Mb) or even about eight-fold in comparison to mammals (~3000 Mb). Interestingly, retronuons themselves generate this genome inflation and thus blunt their own 'virulence' in analogy to strategies of germs that reduce their virulence as mentioned above.

One may draw the following analogy: In a long-drawn trench war enemy artillerists (retroelements) infiltrate the opponent's field fortifications (host genome) and shoot cannons straight into the air (retroposition). Initially, the carnage inflicted by the projectiles is immense. Let's assume, however, that the remaining resident soldiers (genes, gene segments, regulatory

elements) are still viable and may move away from the craters (introns/intergenic regions) generated by the projectiles and thus geographically expand the fortifications (genome inflation). The next rounds of artillery will be less damaging until the soldiers are so widely separated that future rounds of artillery inflict very little damage. With time, for example, the craters fill with rainwater and serve as reservoirs for the maintenance of the resident forces (exaptation). What happens to the infiltrators? Some will also be hit by their own projectiles; they may multiply, but also may be killed captured, tied, by resident forces (host defense) and thus reduced in their effectiveness to operate the cannons. The resident forces may even tolerate them since they do not inflict much damage anymore, but their activities turn out to be useful in one way or the other (recruitment/exaptation). In such a constellation, the resident forces may have an advantage over the opposing forces and win the war (selective advantage) although nobody would have thought and planned so, at the initial stages of battle.

Of course, an opposite strategy that may be equally effective in banning retrons from taking a foothold in genomes is tightening genomes. Organisms with very few and small introns and very little space between genes/operons can select efficiently against the spread of retrons, and a potential parasite or symbiont can only persist when a certain 'consideration' develops towards the host. It is interesting to note that the expansive strategy is mostly associated with complex multicellular organisms with long generation times, while the contractive strategy is common in relatively simple organisms with very short generation times.

Does reverse transcriptase have a functional role in a modern cell, or do retroelements keep the gene alive?

A prerequisite for a full comprehension of retroposition is understanding the evolution and function of the enzyme reverse transcriptase in the cell and during biogenesis of retroelements (Becker, 1996; Doolittle & Feng, 1992). A fundamental question remains as to why reverse transcriptase still is present in so many lineages? Reverse transcriptase, the key enzyme for the transition from RNA to DNA as genetic material may have arisen from an RNA replicase during the final stages of the RNP world (Darnell & Doolittle,

1986; Flavell, 1995; Brosius, 1999c). It is interesting to note that a process that had its heyday several billion years ago during transition from the RNP to the DNA/RNA/protein world is still very active today doing the same as before – converting RNA into genomic DNA – as if the process had a beginning but in certain lineages no end (Jurka, 1998). Rather than a noise from bygone eras, reverse transcription has a dramatic effect on genomes of certain lineages even in quite recent times. If the situation of the almost completed sequence of human chromosome 22 is representative for the entire human genome, almost 40% of the DNA corresponds to retrons (Dunham et al., 1999). These 40% are merely the retrons that we can detect with our current biomathematical analyses of genomes. It is quite conceivable, that almost all of the remaining 55–57% of the genome that does not encode exons is derived from TEs, mainly retrons. Apart from its historical role in converting RNA-based to DNA-based genomes and playing an instrumental role in gigantic genome expansions in certain lineages, an additional function of reverse transcriptase in modern cells remains obscure. In 1985, David Baltimore summarized it as follows: "... there has been no experimental demonstration that reverse transcription has an obligatory role in normal cellular physiology" (Baltimore, 1985). This is still true one and a half decades later, with the noted exception of a specialized reverse transcriptase activity in the maintenance of telomeres (Blackburn, 1992). It is unlikely, however, that the enzymes involved play a role during generation of other retrons (but see (Pardue et al., 1996) where several retroelements were exapted as telomeres in *Drosophila*). Solving the question whether a cellular reverse transcriptase is involved in some vital cellular activity will be extremely challenging. Numerous copies of incapacitated and especially intact and active genes encoding reverse transcriptase provided by retroelements such as proviruses and LINE elements virtually exclude an experimental approach at this juncture. It is accepted that retroelements are very ancient; some authors state that "it seems plausible that retroelements were present at the genesis of living systems" (Flavell, 1995). Others "propose that the occasion of the juxtaposition (of a reverse transcriptase and a prokaryotic transposase) coincided with the invasion of primitive eukaryotes by endosymbiotic prokaryotes 1.5–2.0 billion years ago" (Doolittle & Feng, 1992) and "... the LINEs family of transposable elements is phylogenetically more primitive than the LTR-containing elements as reflected in their

biological distribution, the latter not yet having been identified in any protists. . . we can place the origin of the LINEs family after the introduction of organelles into eukaryotes and anticipate that they will not be found in those primitive eukaryotes like giardia. . .” (Doolittle & Feng, 1992). No matter when exactly retroelements arose, it is even plausible that retroelements alone kept reverse transcriptase alive by evading genomic purges (Jordan, Matyunina & McDonald, 1999) and even re-entering a lineage when countermeasures subside. The alternative is still a currently evasive intrinsic function in the cell, where reverse transcriptase is likely to be encoded by a cellular non-retroelement-derived gene. In the former case, retroposition is forced upon the cell by persistent albeit migrant retroelements, nevertheless conveying long-term benefits to a population or species. In the latter case, it is a by-product of an intrinsic function of the cell. Completely sequenced genomes of model organisms may help identify endogenous reverse transcriptase not derived from retroelements by biomathematical analysis. Experimentally, the feat of removing all retroelement-derived functional copies of the enzyme and then testing for remaining retropositional activity and/or loss of function will remain unattainable for some time, owing to their profusion in mammalian genomes.

Why is there so much extra DNA in many genomes even though its future use is not foreseeable?

Evolution cannot plan for the future: “Evolution has no foresight, and a genetic element cannot be selected because it might someday be of help. Once it is there, however, whatever the reason for its presence, such a structure might prove ‘useful’ and then become the target of some selective pressure on the host phenotype” (Jacob, 1982). Hence, it is difficult to understand why features that may have a potential use in the future convey a selective advantage to the host – especially if one considers that these elements may have serious negative impacts. An insightful and timely treatment of these issues can be read in an article by Doolittle (1989). The author concludes as follows: “Why do so many more species have repetitive sequences, or introns, or excess DNA in general, than we would otherwise, on the basis of our adaptationist understanding of molecular biology, have expected? I submit that an answer lies in the statement that species that have these things more frequently speciate than those which do

not, regardless whether one regards this as a statement about selection as a process or selection as a force. Equally, one might argue that clades whose species retain relatively many of these genomic components less frequently go extinct for failure to adapt to environmental change than do clades whose species do not, even though there is no meaningful sense in which individual organisms within species of the first sort of clade are ‘better adapted’ at the organismal level than are individual organisms within species of clades of the second sort.” In other words, TEs including retrons are found in many extant species because many species that purged themselves of TEs for some short- to mid-term benefit (not being able to ‘know’ about their future exaptive use) are now extinct.

While the dream of solving many questions concerning the history and functional significance of retroposition will largely come true due to the advent of comparative (total) genomics, some questions may remain elusive forever, as intact DNA from long extinct lineages will likely be lost for all time.

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Appendix

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