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## Research

# Extensive somatic L1 retrotransposition in colorectal tumors

Szilvia Solyom,<sup>1,10</sup> Adam D. Ewing,<sup>2,10</sup> Eric P. Rahrman,<sup>3</sup> Tara Doucet,<sup>1,4</sup> Heather H. Nelson,<sup>5</sup> Michael B. Burns,<sup>3</sup> Reuben S. Harris,<sup>3</sup> David F. Sigmon,<sup>1</sup> Alex Casella,<sup>1</sup> Bracha Erlanger,<sup>6</sup> Sarah Wheelan,<sup>6</sup> Kyle R. Upton,<sup>7</sup> Ruchi Shukla,<sup>8</sup> Geoffrey J. Faulkner,<sup>7,8,9</sup> David A. Largaespada,<sup>3</sup> and Haig H. Kazazian, Jr.<sup>1,11</sup>

<sup>1</sup>*McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205, USA;*

<sup>2</sup>*Center for Biomolecular Science and Engineering, University of California at Santa Cruz, Santa Cruz, California 95064, USA;*

<sup>3</sup>*Department of Genetics, Cell Biology and Development and Pediatrics, Masonic Cancer Center, University of Minnesota, Minneapolis, Minnesota 55455, USA;* <sup>4</sup>*Pre-doctoral training program in Human Genetics, McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205, USA;* <sup>5</sup>*Division of Epidemiology and Community Health, Masonic Cancer Center, University of Minnesota, Minneapolis, Minnesota 55455, USA;* <sup>6</sup>*Department of Statistics and Oncology, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205, USA;* <sup>7</sup>*Cancer Biology Program, Mater Medical Research Institute, South Brisbane, Queensland 4101, Australia;* <sup>8</sup>*Division of Genetics and Genomics, The Roslin Institute and Royal (Dick) School of Veterinary Studies, University of Edinburgh, Easter Bush, EH25 9RG, United Kingdom;* <sup>9</sup>*School of Biomedical Sciences, University of Queensland, Brisbane, Queensland 4072, Australia*

L1 retrotransposons comprise 17% of the human genome and are its only autonomous mobile elements. Although L1-induced insertional mutagenesis causes Mendelian disease, their mutagenic load in cancer has been elusive. Using L1-targeted resequencing of 16 colorectal tumor and matched normal DNAs, we found that certain cancers were excessively mutagenized by human-specific L1s, while no verifiable insertions were present in normal tissues. We confirmed de novo L1 insertions in malignancy by both validating and sequencing 69/107 tumor-specific insertions and retrieving both 5' and 3' junctions for 35. In contrast to germline polymorphic L1s, all insertions were severely 5' truncated. Validated insertion numbers varied from up to 17 in some tumors to none in three others, and correlated with the age of the patients. Numerous genes with a role in tumorigenesis were targeted, including *ODZ3*, *ROBO2*, *PTPRM*, *PCMI*, and *CDH11*. Thus, somatic retrotransposition may play an etiologic role in colorectal cancer.

[Supplemental material is available for this article.]

Over two-thirds of our genome may stem from “jumping genes” (de Koning et al. 2011). Three classes of retroelements are known to be currently active and a source of human disease: long interspersed elements (LINEs), the prototype of which is the RNA polymerase II transcribed L1; short interspersed elements (SINEs), consisting essentially of RNA polymerase III transcribed *Alus*; and SVAs (SINE-R/VNTR/*Alus*) that are intermediate in size relative to *Alus* and L1s, and are likely transcribed by RNA polymerase II. A fourth class of retroelements in our genome, human endogenous retroviruses (HERVs) is considered immobile. Full-length L1s are not only responsible for mobilizing themselves, but also for mobilizing the nonautonomous *Alu* (Dewannieux et al. 2003) and SVA retrotransposons (Ostertag et al. 2003; Hancks et al. 2011; Raiz et al. 2011), inactive L1s (Moran et al. 1996), small RNAs (Gilbert et al. 2005), and classical mRNAs, thereby creating processed pseudogenes (Esnault et al. 2000; Wei et al. 2001; Ohshima et al. 2003).

Although there are about half a million L1s in the human genome, only the human-specific L1s (L1Hs) are currently active, represented in each individual by about 800 germline copies

(Ewing and Kazazian 2010), including ~200 full-length sequences (Boissinot et al. 2000). According to conservative estimates there are only about 100 active L1Hs in any human diploid genome that are retrotranspositionally competent, of which six from the reference genome and 37 from six other genomes are known to be highly active (“hot”) (Brouha et al. 2003; Beck et al. 2010). L1s retrotranspose through a process called target-primed reverse transcription (TPRT) (Luan et al. 1993; Cost et al. 2002) with the help of the L1-encoded proteins open reading frame 1 protein (ORF1p) and ORF2p. Endonuclease and reverse transcriptase activities for L1 integration are provided by ORF2p (Mathias et al. 1991; Feng et al. 1996). The hallmarks of TPRT are the addition of a new poly(A) tail to the integrated sequence and target-site duplication (TSD), usually 6–20 bp in length. A fraction of retrotransposition events are also associated with 3' transduction, the comobilization of 3' flanking DNA sequences (Holmes et al. 1994; Moran et al. 1999; Goodier et al. 2000; Pickeral et al. 2000), resulting from transcriptional read-through of the weak L1 poly(A) signal and preferential use of a stronger downstream poly(A) signal. Most de novo L1 retrotransposition events are 5' truncated (Gilbert et al. 2005), with one extreme truncation described where the whole L1 sequence was missing and only the 3' transduced sequence was present (Solyom et al. 2012).

Active mobile elements are not only a significant source of intra- and interindividual variation, but can also act as insertional

<sup>10</sup>These authors contributed equally to this work.

<sup>11</sup>Corresponding author

E-mail [kazazian@jhmi.edu](mailto:kazazian@jhmi.edu)

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mutagens. There are 97 known disease-associated retrotransposon insertions into protein-coding genes (Hancks and Kazazian 2012; van der Klift et al. 2012), which is an underestimate, as conventional mutation screening methods are not designed to amplify large insertions. Of these nearly 100 cases, 25 are caused by L1s, 60 by *Alus*, eight by SVAs, and four by poly(A) sequence originating from an unidentifiable source (Hancks and Kazazian 2012; van der Klift et al. 2012). Of these insertions, 30 occur in cancer cases, including four in colon cancer patients (Miki et al. 1992; Su et al. 2000; Kloor et al. 2004; van der Klift et al. 2012). While three of the four colon cancer cases involve predicted germline or early somatic insertions, a somatic L1 insertion occurred in the *APC* gene in colon cancer (Miki et al. 1992).

In addition to acting as insertional mutagens, retrotransposons can disrupt gene function and genomic integrity in many other ways. These include recombination-mediated gene rearrangements, genetic instability, transcriptional interference, alternative splicing, gene breaking, epigenetic effects, the generation of DNA double-strand breaks, and the expression of small noncoding RNAs (for review, see Goodier and Kazazian 2008; Beck et al. 2011). All of these mechanisms are compatible with a tumorigenic potential of these elements. Retrotransposon overdose is another potential scenario in malignancy and could result in increased insertional mutagenesis, toxicity, or other oncogenic effects. Indeed, the overexpression of L1 ORF1p was observed in certain tumors (Bratthauer and Fanning 1992; Asch et al. 1996; Su et al. 2007; Harris et al. 2010), and RNAi-mediated silencing of L1s resulted in reduced proliferation and differentiation of tumorigenic cell lines (Oricchio et al. 2007). In addition, overexpression of *Alu* elements may exert disease through RNA toxicity (Kaneko et al. 2011). Thus, the cell likely has intrinsic defense mechanisms to prevent retrotransposon overexpression, including methylation (Yoder et al. 1997; Bourc'his and Bestor 2004) and the expression of several host proteins, such as APOBEC3 family members (Bogerd et al. 2006; Chen et al. 2006; Muckenfuss et al. 2006; Stenglein and Harris 2006) or DNA repair enzymes (Gasior et al. 2006; Suzuki et al. 2009; Coufal et al. 2011).

Here we applied two high-throughput L1-targeted resequencing methods to discover retrotransposon activity in colorectal cancers. We identified numerous nonreference L1 insertions not present in paired normal tissue and report a high retrotransposon insertion rate in tumors. We characterized insertion size and TSDs in cancer tissue, confirming that L1s primarily mobilize in cancer via TPRT. The data suggest the importance of retrotransposition in the biology of colorectal tumorigenesis.

## Results

### L1 display through high-throughput sequencing

We applied two next generation resequencing methods—hemispecific PCR coupled to Illumina sequencing (L1-seq) (Ewing and Kazazian 2010) and retrotransposon capture sequencing (RC-seq) (Baillie et al. 2011) to interrogate the retroelement load of colorectal tumors. Approximately 800 nonreference L1Hs copies had been located from individual blood or lymphoblastoid cell lines by L1-seq—the same number as represented by the hg18 reference genome assembly, indicating its capacity to recover essentially all germline L1Hs elements (Ewing and Kazazian 2010). Here, we applied this method to recover somatic insertions from malignant tissues. RC-seq has previously been used to identify somatic mosaicism associated with L1, *Alu*, and SVA mobilization in the brain

(Baillie et al. 2011). Its use of sequence capture for retrotransposon enrichment contrasts with the use of PCR by L1-seq; as a result, RC-seq is expected to cover a broader range of insertions, but with less depth per insertion than L1-seq. A highly multiplexed version of RC-seq was applied to assess whether somatic L1Hs insertions were identified by both approaches.

We sequenced DNA from 16 colorectal tumors and matched normal colons using a pooled L1-seq-based approach. The 16 tumor/normal pairs (32 samples total) were separated into four libraries of eight samples each denoted “colo1/tumor,” “colo1/normal,” “colo2/tumor,” and “colo2/normal.” We sequenced one lane for each library on an Illumina HiSeq 2000 instrument with the exception of colo1/normal, where two lanes of data were generated. The total number of reads generated for each library can be found in Supplemental Table S1.

Using computational methods outlined in Ewing and Kazazian (2010), we identified clusters of reads localized 3' of predicted insertion sites. We required 100 reads spanning at least 100 bp (“high stringency”) as a minimum for L1 detection, which yields a specificity of >90% based on recovery of reference L1 insertions and nonreference sites discovered in previous studies (see Supplemental Figs. S1, S2 for an exploration of cutoff parameters). Using these criteria, we identified 764 reference L1Hs insertion sites present in NCBI36/hg18 and 400 nonreference insertion sites from the colo1 data. From the colo2 data we identified 816 reference and 433 nonreference insertions. Combining the data, we found 819 reference L1Hs elements and 635 nonreference elements, 336 of which had not been previously cataloged. Many of these uncataloged elements are new somatic insertions in the tumor. In total, 38% of reference and 35% of nonreference insertions were in gene annotations based on UCSC Known Genes. The distribution of L1 insertions detected by L1-seq in this study is shown in Figure 1.

Our primary interest in generating these data was in finding insertions present either in a cancer pooled library or in a normal



**Figure 1.** Genomic distribution of L1 insertions. Outer rings show the density of detected insertion sites for reference (gray) and nonreference (black) L1s. The approximate locations of the 72 PCR-validated somatic insertions are indicated by dots inside the circle. Note that 69 of the 72 insertions were successfully sequenced.

pooled library and not present in the corresponding paired normal or tumor library. Turning to these, with the same stringency cutoffs as above, we found 35 putative insertions only in colo1/tumor, four only in colo1/normal, 50 predictions only in colo2/tumor, and eight only in colo2/normal. Decreasing the requirements for predicted insertions to 10 reads spanning at least 100 bp ("low stringency"), we found 69 potential insertions only in colo1/tumor, 173 only in colo1/normal, 75 only in colo2/tumor, and 42 only in colo2/normal. The dramatic increase in predictions for colo1/normal only with decreasing stringency is an effect of the higher coverage in the colo1/normal versus colo1/tumor (Supplemental Table S1), as two lanes of sequence were generated for colo1/normal.

Five of the L1-sequenced colorectal tissue pairs were bar-coded, pooled, and analyzed by shallow, multiplexed RC-seq (10 libraries, ~75 million paired-end Illumina GAIIx reads). A total of 26,903 nonreference genomic insertions were detected by at least one read (Supplemental Table S2). Of these, 358 were (1) found in only one donor, (2) were not identified in RC-seq previously performed on pooled blood (Baillie et al. 2011) or databases of retrotransposon polymorphisms (Huang et al. 2010; Iskow et al. 2010; Ewing and Kazazian 2011), and (3) could be annotated with high confidence due to detection by multiple unique amplicons. Of this set, 96 were only found in tumor, including eight L1, 83 *Alu*, and five SVA. A total of 39 insertions were found only in nontumor samples and 223 were found in both tumor and nontumor. The tumor:nontumor ratios for L1, *Alu* and SVA, were ~8:1, 2.5:1, and 2:1, respectively. *AluY* and L1-Ta/pre-Ta were detected, but no HERVs were detected.

### PCR validation

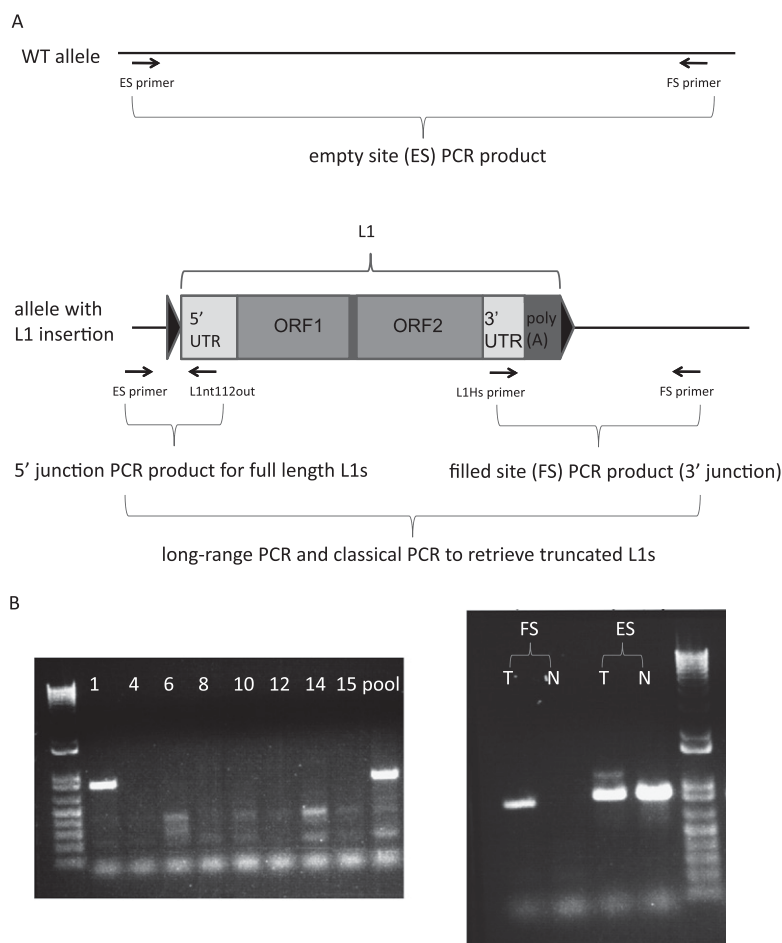
We applied a step-wise PCR amplification scheme to validate insertion sites from L1-seq data and to determine both 5' and 3' junctions of L1Hs elements identified by L1-seq. Primary validations focused on confirming the presence of de novo L1 inserts by amplifying their 3' junction and determining which of the eight patients carried the insertion within a DNA pool (PCR scheme and primer design performed according to Ewing and Kazazian 2010) (Fig. 2A,B). An L1 insertion was considered to be validated as tumor specific if the filled site (L1-containing) PCR product was present in the tumor, but not in the paired normal tissue, and in the case of heterozygous autosomal insertions, the empty site PCR product (WT allele) was amplified from both members of the tissue pair. Using a single PCR condition to amplify the 3' junctions, we PCR-validated 26/40 and 37/51 insertions from the colo1 and colo2 high-stringency data sets, respectively. We also set out to PCR amplify 16 colo1 insertions represented by less than 100 Illumina reads

and were able to validate nine. Thus, we PCR-validated before sequencing the 3' ends of 72 of 107 putative insertions (Supplemental Table S3; Supplemental Text S1).

Interestingly, among 12 high-stringency putative insertions from normal colon of the combined colo1 and colo2 data sets, none could be validated. Possible explanations for false positives in the L1-seq data include PCR artifacts arising during library preparation, suboptimal PCR conditions used for validation, or L1 insertion into repetitive sequences, refractory to successful primer design.

Our stepwise PCR amplification scheme continued by retrieving the 5' junctions of tumor-specific L1 insertion events. Several empty-site PCRs had already yielded a higher molecular weight band exclusively in the tumor, which in each case was verified to be a highly truncated L1 element (Fig. 2B). In the remaining cases, long-range PCR and a PCR specifically designed to amplify the 5' end of a full-length L1 were used to retrieve the 5' junction.

Altogether, out of 72 cases where the insertions were PCR-validated to be tumor specific, we successfully sequenced either the 3' or the 5' junction in 69 cases (Supplemental Table S3; Supple-



**Figure 2.** PCR validation scheme of L1-seq results. (A) The three-step PCR validation scheme and location of primers used. Triangles symbolize TSD. (B) PCR validation of the 3' junction (ins. 7). This insertion is in tumor 1 of the eight DNA samples that had been pooled for Illumina sequence analysis (left), while the right panel shows it is present exclusively in the tumor, but not in the normal colon. The higher molecular weight band visible above the ins. 7 empty site PCR product in the tumor is a highly truncated L1. (T) Tumor; (N) normal colon; (FS) filled site PCR product; (ES) empty site PCR product.

mental Text S1). For 35 insertions, we sequenced both junctions, enabling us to characterize TSDs and L1 insertion size in cancer tissue (Table 1). Surprisingly, all of the tumor-specific insertions were highly truncated, the mean L1 insertion size being 585 bp, excluding the poly(A) tail (Table 1).

Using a PCR designed to amplify full-length L1 insertions, we failed to amplify the 5' end of any of the remaining tumor-specific insertions where 5' junctions could not be identified with the previous PCR approaches. On the other hand, three of 10 germline polymorphic insertions had an intact 5' end, in agreement with 30% of reference L1 elements being full length (Pavlicek et al. 2002). This difference between full-length L1 insertions in tumors (~0%) versus full-length insertions among polymorphic germline L1s (~30%) is statistically significant ( $P = 0.016$ , Fisher's exact test) and is a clear departure from what is observed from the reference genome and from heritable nonreference insertions (Ewing and Kazazian 2010).

L1-seq results on five tumors were corroborated by RC-seq. Eleven high-confidence L1s hits were found in these cancers by L1-seq, out of which four were also detected by RC-seq at either the 5' or 3' junction (ins. 5, 9, 14, and 32) (Supplemental Tables S2, S3; Supplemental Text S1). Among eight high-confidence L1 insertions within genes detected by RC-seq, but missed by L1-seq, one was validated as present in tumor 10, targeting the *DGKI* gene (Supplemental Table S2) (Fig. 3). Of the remaining putative tumor-specific insertions that could be PCR-amplified, six of eight L1s, 30 of 57 *Alus*, and six of 11 SVAs were present in both tumor and paired normal tissue. No other confirmed tumor-specific insertions from the L1-seq data were found by RC-seq.

In order to determine whether L1s are frequently mobilized in other nonmalignant somatic tissues, we performed L1-seq on genomic DNA extracted from cerebrum, liver, and testis samples from two other individuals (cadaver samples) who had died from arteriosclerotic cardiovascular disease. None of the Illumina high-stringency sequence peaks suggestive of somatic L1 insertion could be PCR-validated, implying that the high rate of somatic L1 insertions observed in colon cancer was specific to the malignant tissue. Thus, we have no evidence from our data of somatic insertions in normal colon, liver, testis, or cerebrum.

### Characterization of tumor-specific insertions

Intriguingly, the number of validated L1 insertions varied widely from tumor to tumor with up to 17 insertions in some and none in three others (Fig. 3). Most retrotransposition events showed hallmarks of TPRT, namely TSD (27/35), L1 endonuclease cleavage site, the presence of L1 poly(A) tail, frequent 5' inversion (10/35), and in one case, a 3' transduction (Table 1). However, we note that a substantial fraction of these somatic insertions (8/35) lacked a TSD and six of these lacked a discernible endonuclease cleavage site, suggesting that they were endonuclease-independent insertions (Morrish et al. 2002). Two insertions ("3" and "21") contained 3' sequence from other chromosomes, but lacked a poly(A) tail in between the two sequences that would indicate a 3' transduction. Thus, they are likely cancer-associated recombination events (Supplemental Table S3; Supplemental Text S1).

Numerous genes were targets for insertional mutagenesis in colon tumors by L1s that are represented in the COSMIC database (Catalogue of Somatic Mutations in Cancer, <http://www.sanger.ac.uk/genetics/CGP/cosmic/>). Examples include *PTPRM* (protein tyrosine phosphatase, receptor type, M), *ODZ3* (odd Oz/ten-m homolog 3), *ROBO2* (roundabout, axon guidance receptor, ho-

molog 2), *PCMI* (pericentriolar material 1), and *CDH11* (cadherin-11). *PCMI* and *CDH11* are also represented in Sanger's Cancer Gene Census (<http://www.sanger.ac.uk/genetics/CGP/Census/>). Interestingly, according to COSMIC, in the large intestine these genes were mutated with the following high frequencies: *PTPRM* (50%), *ODZ3* (100%), *ROBO2* (15%), *PCMI* (12%), *CDH11* (52%). All of our hits were intronic and were PCR validated as well as sequenced. Additional interesting genes with a potential role in malignancy were also targeted, for instance, *RUNX1T1* (runt-related transcription factor 1), a member of the myeloid translocation genes. Interestingly, somatic *RUNX1T1* point mutations were not only found in colorectal cancers (Wood et al. 2007), but the product of the related *RUNX3* gene regulates L1 expression (Yang et al. 2003).

### Cellular timing of L1 retrotransposition

In an effort to determine at what point in tumorigenesis the L1 insertions occurred, we developed three lines of evidence: analysis of SNPs in sequence flanking the L1 insertion and the empty site, the number of empty site X chromosome alleles in males who had an L1 insertion into the X, and the presence/absence of the L1 insertion in a second section of a particular tumor in which an L1 insertion occurred.

First, we found three insertions (C2, C4, and insertion 31) with flanking heterozygous SNPs. For C2 (*ODZ3* gene) there was one SNP, for C4 there were four SNPs, and for insertion 31 there was one flanking SNP (Supplemental Fig. S3A). The presence of both alleles of the particular SNP in the empty site chromosomes is informative in case of no aneuploidy at the respective alleles. If the insertion occurred at the initiation of the tumor (the one-cell stage), the filled site would contain one allele and the empty site would contain the other allele only. If both alleles are present in the empty-site chromosomes, then the insertion likely occurred after the one-cell stage of the tumor. The data showed that all six SNPs near three different insertions in the empty-site chromosomes were heterozygous, suggesting that the insertions occurred after the initiation of tumorigenesis. We carried out array comparative genomic hybridization (aCGH) and found no copy-number gain or loss in the chromosomal arm of these SNPs (data not shown), although small chromosomal aberrations at the respective alleles cannot be ruled out.

Second, we found insertions (D9, D12, and E3) into the X chromosome in two males. If the insertion occurred at the one-cell stage and the male did not have X-chromosome aneuploidy, the tumor should lack an empty site. However, in all cases, we found an empty-site band by PCR, indicating again that the insertions occurred after the one-cell stage of tumorigenesis (Supplemental Fig. S3B contains data on D9 and D12). Again, aCGH showed that both males had a single X chromosome.

Third, we obtained a second portion of tissue from a number of tumors and determined whether the insertions found in the first tumor tissues could be confirmed in the second tumor sample. In three of seven instances (insertions C2, D7, and E3) we were able to confirm the insertion in a second tumor sample, suggesting a relatively early event in tumorigenesis. Four other insertions ("31," "A10," "C4," "D12") were present in the first tumor section, but not in the second (Supplemental Fig. S3C). In tumor 2853, two insertions were studied: E3 was present in both tumor portions, while D12 was not, suggesting that these two insertions occurred at different times and in different cells of the tumor (data not shown). Furthermore, heterozygosity for L1 flanking SNPs in in-

**Table 1.** L1Hs insertions in cancer with both 5' and 3' junctions characterized

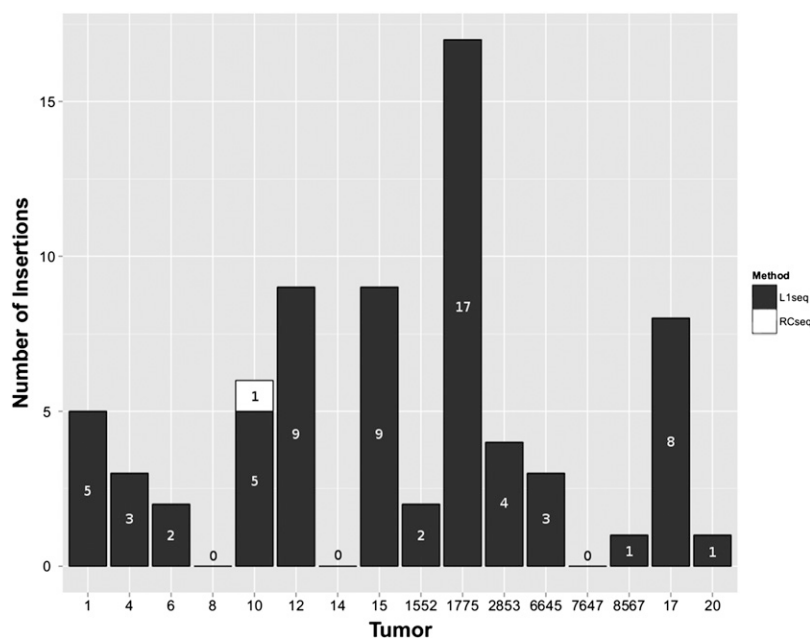
Insertion	Tumor	Chr	Chr location (hg18)	Chr location (hg19)	L1 5' junction	Inferred L1 size (bp)	Identity to LIRP (%)	Poly(A) length (bp)	TSD	Endo site	Notes
Ins. 5	1	10	56731539–56731701	57061533–57061695	5898–5934	122	100	73	AACAGAAGTCTCA (13 bp)	TGTT/GA	5' Junction with RC-seq
Ins. 6	15	11	20389184–20389665	20432608–20433089	5779–5432 5780–5965 5786–5893	588	100	17	AAATACCTTTAGTT (14 bp)	ATTT/TA	Partly inverted L1
Ins. 7	1	11	36145260–36145460	36188684–36188884	5786–5893	234	100	16–70	GAAGAAATATC (11 bp)	CTTC/AG	
Ins. 9	1	11	114813023–114813333	115307813–115308123	5161–5849	859	100	71	A (1 bp)	T/AA	3' Junction also with RC-seq
Ins. 24	10	3	1873003–1873267	1898003–1898267	5473–6019	547	99	32–37	AGAAITTA (8 bp)	TTCT/AA	Potentially endo-independent
Ins. 30	15	3	175764832–175764906	174282138–174282212	5762–6019	258	99	42–43	None	TTTC/AG	3' Junction also with RC-seq
Ins. 32	1	4	166273011–166273097	166053561–166053647	5399–6019	621	100	39	None	TTAT/CA	Potentially endo-independent
Ins. 35	1	5	165976534–165976920	166043956–166044342	5640–6019	380	100	23–104	None, 6-bp deletion	GTTT/TT	Potentially endo-independent
Ins. 36	6	6	91768021–91768340	91711300–91711619	5818–6018/6019	202	99	26–28	(T)AGTTGGTACG TTTTT (15/16 bp)	AACT/AA	Inverted poly(A) inserted in front of L1 nt. 5818; untemplated nt.-s AAAGCCTG inbetween
Ins. 47	15	11	90728127–90728299	91088479–91088651	5693–5894	327	100	74	None	ACAT/TG	Potentially endo-independent
Ins. 50	10	16	63628221–63628561	65070720–65071060	5245–5172	775	100	21 + 18–29 (3' transduction)	AAGAACTA (9 bp)	TCTT/AG	Insertion into CDH11; L1 inversion; 124-bp 3' transduction from chr 1
Ins. 57	4	7	54612665–54612869	54645171–54645375	5308–4921 5331–5628 5626–5916	1077	100	83	AAGAAATACCATA GGTTGG (18 bp)	TCTT/AT	Partly inverted L1 with L1 deletion
Ins. A2	6645	1	118179594–118179855	118378071–118378332	5626–5916	394	99	45	AAAAATTAC AACTACA (17 bp)	TTTT/AC	Untemplated TCC nt.-s at 5' breakpoint
Ins. A4	1775	1	186265021–186265327	187998398–187998704	5118–5800	902	99	17	None	ATAT/TG	Potentially endo-independent
Ins. A5	1775	10	52852490–52852780	53182484–53182774	5560–5890	460	N/A	15	AAAAACAAA AAA (13 bp)	TTTT/AA	Untemplated AA nt.-s at 5' breakpoint
Ins. A8	17	12	18980506–18980824	19089239–19089557	5498–6019	522	99	46	AAAAAG (7 bp)	TTTT/AT	
Ins. B2	1552	17	13914476–13914781	13973751–13974056	5832–6019	188	100	25–34	GAAT (6 bp)	TTTC/AT	
Ins. B4	1775	18	67449071–67449391	69298091–69298411	5525–5885	495	100	21	AAAACTCC (8 bp)	AAAT/AC	
Ins. B5	17	2	140975700–140975940	141259230–141259470	5695–5907	325	100	35	AAAAGTAGC AAAT (13 bp)	TTTT/GA	
Ins. B6	1775	2	188158976–188159118	188450731–188450873	4909–5371	1111	N/A	11	AAAAATGCATAA (12 bp)	TTTT/AT	
Ins. B7	1775	2	188578469–188578710	188870224–188870465	5570–5907	450	100	31	TAAAAGATCTTA AATA (16 bp)	TTTA/AA	
Ins. B9	1552	3	141880688–141880901	140397998–140398211	3161–2454 4510–4608	2218	99	27	AAAAAGTTA CAGGTATT (16 bp)	TTTT/AA	Partly inverted L1 with L1 deletion
Ins. B10	1775	4	93068869–93069099	92849846–92850076	5896–6019	124	99	36–40	AAAG (4 bp)	CTTT/AC	

(continued)

**Table 1.** *Continued*

Insertion	Tumor	Chr	Chr location (hg18)	Chr location (hg19)	L1 5' junction	Inferred L1 size (bp)	Identity to L1P (%)	Poly(A) length (bp)	TSD	Endo site	Notes
Ins. C4	6645	5	62265644–62265857	62229888–62230101	5270–5750	750	99	26	None	TTTA/TG	Potentially endo-independent
Ins. C6	1775	6	87790188–87790349	87733469–87733630	5807–5903	213	100	16	AAGAGATTGG CAAAATCA (17 bp)	TCTT/AT	
Ins. C7	1775	6	132427126–132427288	132385433–132385595	5703–5319 5721–5891	384	100	28	AAAAGAAA TATGTC (14 bp)	TTTT/AG	Partly inverted L1 with L1 deletion
Ins. C8	1775	7	43206340–43206484	43239815–43239959	5892–6019	128	98	27–50	None, 1-bp deletion	CGTC/AT	Potentially endo-independent
Ins. C10	6645	7	71370938–71371265	71733002–71733329	5830–6019	190	100	48–49	A (1 bp)	T/AA	
Ins. D5	1775	8	111653589–111653790	111584413–111584614	4861–4525 4922–5269	1435	100	40	A (1 bp)	T/CA	Partly inverted L1 with L1 deletion
Ins. D7	1775	8	140519447–140519608	140450265–140450426	5214–5867	806	N/A	30	AAAAAAGGACT (11 bp)	TTTT/AT	
Ins. D9	17	X	18694952–18695143	18785031–18785222	4854–5680	1166	99	46	ATAAAAATGAG (11 bp)	TTAT/AG	
Ins. D10	1775	X	83860994–83861183	83974338–83974527	4504–4011	1516	N/A	76	TAAACACAG AGAACA (14 bp)	TTTA/AT	Partly inverted L1
Ins. D11	1775	X	87182336–87182515	87295680–87295859	5913–6019	107	100	21–26	None, 1-bp deletion	CTTT/AT	
Ins. D12	2853	X	108163640–108163825	108276984–108277169	5399–5361 5549–5908	510	100	59	AAAAAGTTAA CTTGTT (15 bp)	TTTT/AT	Partly inverted L1 with L1 deletion; untemplated "A" in between them
Ins. E1	1775	X	108585628–108585763	108698972–108699107	5655–5644 5669–5890	363	100	13	AAAATCAACA TACCCA (17 bp)	TTTT/AA	Partly inverted L1 with L1 deletion

The L1 5' junction nucleotide position extends until the point where the sequence chromatogram was readable. Inferred L1 size is calculated by hypothesizing that no further L1 deletions occurred. Poly(A) length cannot be determined unambiguously by Sanger sequencing due to different levels of polymerase slippage from the forward and reverse sequencing directions. Ins. 5–57 are *colo1*, and ins. A2–E1 are *colo2* L1s.



**Figure 3.** Distribution of somatic L1 insertions in tumors. Insertions in black were detected by L1-seq, while the insertion in tumor 10 in white was detected by RC-seq only.

sertions 31 and C4 in the original tumor sample, as well as the absence of the insertion from the second tumor portion both suggest late insertion events. Thus, from the combination of these data on a small sample size we conclude that most, if not all, of the studied L1 insertions occurred after the initiation of the tumor. However, a minority of the insertions may have occurred at an early stage of tumorigenesis. Furthermore, we cannot exclude the possibility of tumor blood vessels or infiltrating lymphocytes as contributing alternative explanations for some of the results.

#### Effects of L1 retrotransposition on measures of cellular instability

To address the increased rate of somatic L1 retrotransposition in tumors, we assessed the genomic landscape of the tissue samples by aCGH, microsatellite instability (MSI), and L1 promoter methylation status. A previous report demonstrated a correlation of genome-wide DNA methylation status of tumors with increased de novo L1 insertions (Iskow et al. 2010). We assessed the methylation status of the L1 promoter at four different CpG sites. Although L1 promoter hypomethylation was found in tumor samples compared with paired normal tissue, no correlation was observed between L1 methylation status and the number of L1 insertions (Fig. 4A).

For aCGH, we analyzed normal and tumor tissue from six patient samples, two that possessed the greatest number of de novo tumor insertions (12 and 1775), two with no tumor insertions (8 and 7647), and the two males mentioned above with both empty and filled insertion sites in X chromosomes (17 and 2853). Tumor samples 8, 12, 17, and 1775 contained complex chromosomal changes, including entire and partial chromosomal gains and losses. Tumor samples 2853 and 7647 presented no detectable aberrations relative to normal tissue. Interestingly, three of the samples (12, 17, and 1775) with complex chromosomal rearrangements had a high number of validated insertions. Likewise,

patient 7647 with no validated L1 insertions had no detectable chromosomal changes. The outliers from the trend of a direct relationship between chromosomal aberrations and L1 insertions were patients 8 and 2853. Interestingly, these latter patients are potentially genetically predisposed to colon cancer due to familial cancer aggregation or a very young age of diagnosis.

In addition to analysis of gross genomic abnormalities, we assessed the status of the mismatch DNA repair pathway by assessing microsatellite expansions in the genome. Seven of the 16 patients were MSI positive. Although two samples with the highest number of somatic L1 insertions were MSI positive, MSI status did not correlate with the number of de novo L1 insertions for each tumor (Fig. 4B).

Interestingly, a statistically significant correlation was observed between the number of insertions and the age of the investigated patients ( $P = 0.01425$ ,  $R^2 = 0.3128$ , where age is the time of surgical sample removal). Eight or more validated insertions were observed only

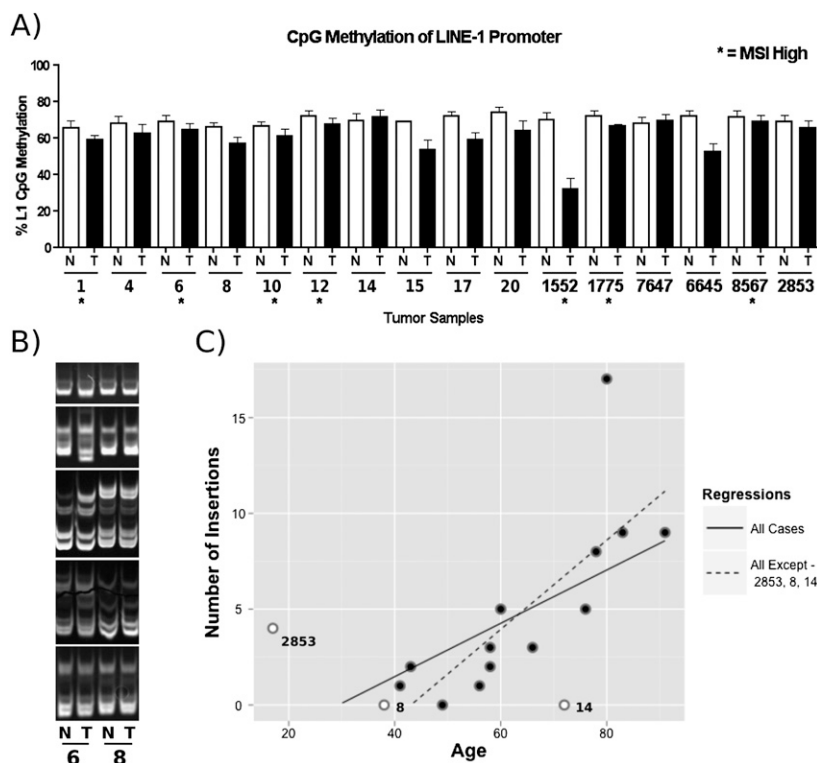
in the tumors of patients 78 yr old or older. An outlier in the correlation was a 72-yr-old with no validated insertions. However, he was the only proband with rectal cancer, but no colon tumor diagnosis. When this patient was excluded from the analysis, as well as cases with a presumed genetic predisposition to colon cancer (familial polyposis case and a 17-yr-old male), an even more significant correlation was observed between the age of sporadic colon cancer patients and L1 activity ( $P = 0.001548$ ,  $R^2 = 0.578$ ) (Fig. 4C).

#### Discussion

Our L1-seq method has revealed a high rate of L1Hs retrotransposition in certain colorectal cancer genomes. The neighboring matched normal colon sample in these 16 cases, as well as cerebrum, liver, and testis from two other individuals yielded no L1 insertions that could be validated, indicating few or no retrotransposition events in these normal tissues.

Iskow et al. (2010) used 454 pyrosequencing to search for de novo L1 insertions in five glioblastomas, five medulloblastomas, as well as leukemia and breast cancer cell lines, but they found no insertions in these cases. However, they identified nine somatic L1 insertions in six of 20 lung tumors. Since TSDs were not reported, the question of whether L1 integration in lung cancer occurs through TPRT remained open.

Here we report that evolutionarily young L1Hs retrotransposons can mobilize themselves through the classical TPRT mechanism in colon cancer genomes at a high frequency. The true retrotransposition rate is likely to be even higher, as our method does not detect insertions mobilized by L1 elements in trans, such as *Alus*, SVAs, most inactive L1s, and processed pseudogenes. In addition, there are likely other L1 insertions in our data set that have not been subjected to validation. Longer tumor-specific 3' transductions will be missed as well, as it is difficult to differentiate between the progenitor and the transduced sequence by their 3' flank with L1-seq.



**Figure 4.** Analysis of factors influencing L1 activity. (A) L1 CpG promoter methylation status performed by quantitative bisulfite PCR analysis. (N) Normal tissue; (T) tumor tissue; (\*) MSI. Replicates of four were done for each data point. (Error bars) Standard deviations. (B) MSI analysis. 6% TBE gel depicting the status of five microsatellite repeats (*BAT25*, *BAT26*, *D2S123*, *D17S250*, *D17346* in descending order) in normal and tumor tissue from two different patients. Tumor tissue "6" contained additional bands and gel shifts compared with the normal tissue, indicating MSI. Samples from "8" demonstrated no differences suggestive of MSI. (C) Correlation of L1 activity with age of the patient at time of surgery. See text for details.

In order to determine why the rate of somatic cell retrotransposition was high in some tumors compared with the normal tissue, we assessed how the following factors correlate with retrotransposition rate: age of colorectal cancer patients, chromosomal aberrations, L1 methylation status, and mismatch DNA repair as reflected by MSI. A clear correlation of retrotransposition activity was observed with the age of colon cancer patients in sporadic cases. Furthermore, we also analyzed genetic instability by aCGH in six samples and found a modest association between the number of chromosomal aberrations and the age of the sporadic colon cancer patients. Altogether, it is possible that the hypomethylated microenvironment of the tumors, together with genetic instability as reflected by MSI and gross chromosomal changes, have a cumulative effect in older patients. Our results are in agreement with the correlation of retrotransposition activity with genome instability during yeast chronological aging (Maxwell et al. 2011).

In this study, we found that genes with a known driver function in cancer are mutagenized by L1Hs elements. The accumulation of retrotransposon sequences is predicted to cause further genetic instability through recombination. Thus, an elevated insertion rate is expected to contribute to tumor evolution. As exemplified by a somatic L1 insertion into the *APC* gene (Miki et al. 1992), it is clear that in some fraction of colorectal cancers retrotransposon insertions can be etiologically significant. Yet, it remains unclear in what fraction of cases retrotransposons initiate malignant transformation and in how many instances they con-

tribute solely to a more aggressive phenotype. SNP, X chromosome, and secondary sampling data from tumor samples suggest that L1 insertions likely occurred at various times after the initiation of the tumor. Although we found no evidence for L1 insertion into colon cancer tumor-suppressor genes or oncogenes that would be indicative of driver mutation-induced tumor clonality, analysis of a larger number of tumors or a deeper sequencing of retrotransposon insertions could uncover such events. We propose that it is possible to estimate insertion timing and tumor heterogeneity more precisely by evaluating pure tumor samples. Our findings are in agreement with a very recent report on retrotransposon insertions in epithelial cancers (Lee et al. 2012). Intriguingly, an intronic L1 integration event was found in that study as well in the *ROBO2* gene in a colon tumor. Additionally, they detected intronic L1 insertions in *CDH12*, while we characterized an insertion into the *CDH11* gene. Thus, the role of cell-adhesion genes in retrotransposon insertion-mediated colorectal tumorigenesis may deserve further investigation.

An unexpected finding of our PCR-based validation is the severely truncated nature of all validated L1 insertions in colon cancer. It was not possible to assess whether this is a general characteristic of the malignant phenotype, of all somatic tissues, or of the gastrointestinal tract in particular, as no de novo L1 insertions could be uncovered from normal colon, liver, testis, and brain. In a transgenic mouse model, 30 of 33 somatic L1 insertions were 5' truncated (Babushok et al. 2006), raising the possibility of a gradual decrease in L1 size from germline to somatic to malignant insertions. In cultured HeLa cells, 94/100 insertions were 5' truncated (Gilbert et al. 2005). This might indicate that some cancer tissues or cultured cells could allow full-length insertions to accumulate. The overexpression of an exogenous L1 element, coupled with the bias toward recovering larger inserts in that assay, and the unknown effects of cell culture conditions on retrotransposition complicate transferring conclusions on the L1 5' truncation rate to cancer tissue. Likewise, it is not understood why the majority of germline L1 insertions are 5' truncated as opposed to *Alu* and *SVA* insertions that are mostly full length, yet also mobilized by L1s (Hancks et al. 2011).

The truncated structure of L1 elements in colorectal cancer may be useful in understanding the mechanism of 5' truncation both in normal and tumor cells. We propose two possible explanations: (1) If TPRT timing is coupled to the cell cycle, the elevated cell division rate of malignant cells may not leave sufficient time to complete integration of long mobile elements; (2) a DNA repair pathway might monitor and remove de novo mobile element insertions in healthy tissues. Once this presumed surveillance pathway is down-regulated in cancer, retrotransposon insertions are not removed efficiently and are allowed to accumulate. At the same time, another or the same DNA repair pathway might specialize in truncating fresh integrants or prohibiting them from

completing retrotransposition. If this process is up-regulated, the truncation rate increases. The efficiency of such a pathway might correlate with insertion size or be sequence specific, thus preferentially targeting L1 elements over *Alu*s and SVAs. Interestingly, nonhomologous end joining (NHEJ) is an important DNA repair pathway candidate with a reported conflicting dual role in regulating retrotransposon insertions, offering an explanation for parallel L1 up-regulation and truncation (Suzuki et al. 2009). We propose that by comparing the genome or transcriptome of tumors with a high rate of retrotransposition to their paired normal tissues, we may discern clues to cellular factors causing L1 mobilization and 5' truncation.

To conclude, the cancerous colon of many patients is the second reported organ beside the brain (Baillie et al. 2011) in which a high rate of retrotransposition occurs. Lung, prostate, and ovarian tumors are also reported to allow a lower level of L1 mobilization (Iskow et al. 2010; Lee et al. 2012), but many other cancer types appear to be nonpermissive for a detectable rate of retrotransposition. All L1 insertions in the colorectal tumors of this study were highly truncated, potentially indicating the footprint of a defective or hyperactive DNA repair pathway in cancer. The cause and effect of retrotransposon mobilization in cancers warrants further investigation.

## Methods

### Human DNA samples

DNA was extracted from human patient tissue samples acquired from the University of Minnesota Tissue Procurement Facility from BioNet (IRB#0805E32181). See Supplemental Table S4 for patient data. Briefly, 2 mg of tissue was digested overnight at 55°C on a rotating platform in 710  $\mu$ L of digest buffer (1 M Tris at pH 8.0, 1 mM EDTA, 1 $\times$  SSC, 1% SDS, 1 Mm NaCl, 10  $\mu$ g/mL Proteinase K). Following digest, DNA was purified using phenol-chloroform-isoamyl alcohol (Life Sciences) isolation protocol.

Human frozen tissue from two Caucasian cadavers with arteriosclerotic cardiovascular disease were obtained from the NICHD Brain and Tissue Bank for Developmental Disorders at the University of Maryland, Baltimore. DNA isolation was done utilizing the AllPrep DNA/RNA Mini Kit (Qiagen).

### Library construction, sequencing, and analysis

#### L1-seq

The library for L1Hs elements was made according to Ewing and Kazazian (2010), while the library for L1, *Alu*, and SVA elements (RC-seq) was constructed according to Baillie et al. (2011). L1Hs elements were TOPO-TA cloned (Invitrogen) and Sanger-sequenced for quality control of the library preparation, and were subsequently sequenced on an Illumina HiSeq 2000 at the Johns Hopkins University Genetic Resources Core Facility High Throughput Sequencing Center.

Pooled L1-seq library sequence data was analyzed as described previously (Ewing and Kazazian 2010), and compared against insertion sites of known reference and nonreference transposable elements (Beck et al. 2010; Ewing and Kazazian 2010, 2011; Huang et al. 2010; Iskow et al. 2010; Witherspoon et al. 2010; Hormozdiari et al. 2011; Stewart et al. 2011). Gene annotations were obtained from UCSC Known Genes (Hsu et al. 2006).

#### RC-seq

The library for retrotransposon capture and sequencing was created utilizing DNA from five pairs of colorectal and normal tissue

(samples 1; 4; 6; 8; 10) by the same method as published previously (Baillie et al. 2011). One significant change was made to the technique, in that liquid phase hybridization was performed as opposed to solid surface, chip-based hybridization. The libraries were then sequenced on an Illumina Genome Analyzer II and aligned to the genome by a computational pipeline that utilized SOAP2 to align reads to the genome and used much the same method as published (Baillie et al. 2011; R Shukla, KR Upton, M Muñoz-Lopez, DJ Gerhardt, JK Baillie, ME Fisher, PM Brennan, A Collino, S Ghisletti, S Sinha, et al., in prep.).

### PCR validation of the Illumina results

A three-step PCR validation protocol was used to validate the next-generation sequencing reads and to retrieve 3' and 5' junctions. As the first step, L1 3' ends together with flanking genomic regions were amplified using the same AC dinucleotide-specific primer of L1Hs as used for Illumina sequencing (L1Hs primer: GGGAGAT ATACCTAATGCTAGATGACAC) and a primer selected from the 3' flanking region based on the reference genome sequence (FS primer). PCR reactions were carried out in 12.5  $\mu$ L of 2 $\times$  GoTaq Green master mix (Promega) in a total volume of 25  $\mu$ L, with 0.8  $\mu$ L of FS primer, 1.5  $\mu$ L of L1Hs primer, and 25 ng of DNA to amplify the filled site. The empty site was amplified with the same conditions, except that 1.5  $\mu$ L of FS primer, 1.5  $\mu$ L of ES primer, and 12.5 ng of DNA were used. Primers were 20 pmol/ $\mu$ L and their location is depicted in Figure 2A. Reactions were incubated for 2 min at 95°C, followed by 30 cycles of 30 sec at 95°C, 30 sec at 57°C, and 1.5 min at 72°C, followed by a final extension of 5 min at 72°C on a PTC-200 Peltier Thermal Cycler. Long-range PCR to recover longer L1 insertions was performed with the Expand Long Template PCR System (Roche) according to the manufacturer's instructions in buffer 1, with 1  $\mu$ L of 20  $\mu$ M FS and ES primers each, and 25 ng of tumor DNA. 5' junctions were PCR amplified using the same conditions as for the 3' junction, except that a primer hybridizing to the L1 5'UTR was used (L1nt112out: GATGAACCCGGTACCT CAGA) together with the respective ES primer, and primer extension time was only 45 sec. FS and ES primer sequences are included in the Supplemental Material (Supplemental Table S1). PCR products were cut out of the gel, extracted with the QIAquick Gel Extraction Kit (Qiagen), and sequenced. See Supplemental Text S1 for Sanger sequence data on insertions.

### Microsatellite instability assays

To assess the MSI status, we utilized five markers recommended by the National Cancer Institute (Bethesda markers): BAT25 and BAT26 to assess mononucleotide repeats (A)<sub>n</sub> and D2S123, D5S346, and D17S250 to assess dinucleotide repeats (CA)<sub>n</sub>. MSI status was determined using previously established protocols (Ashktorab et al. 2003; Muller et al. 2004). Primers were developed by the NCI for screening patients in the clinic.

### L1 methylation status

The methylation level of L1 promoters was performed according to Wilhelm et al. (2010). Briefly, each sample was amplified three times and each amplification was pyrosequenced once. The average of the three was utilized to determine the value of CpG methylation for each of the four positions analyzed for an L1.

### aCGH

DNA from patients 8, 12, 17, 1775, 2853, and 7647 were restriction digested and labeled with fluorochrome Cyanine-5 using random

primers and exo-Klenow fragment DNA polymerase. DNA from a sex-matched control was labeled concurrently with Cyanine-3. The sample and control DNA were combined and array-based comparative genomic hybridization (aCGH) and single nucleotide polymorphism analysis (SNP) was performed with a 180K Cancer CGH+SNP microarray constructed by Agilent Technologies, Inc. that contains ~115,000 distinct biological oligonucleotides and 55,000 SNP sites, spaced at an average interval of 25 KB (for 20,000 cancer-associated CGH probes: one probe/0.5–1 KB). The ratio of sample to control DNA for each oligo was calculated using Feature Extraction software 10.10 (Agilent Technologies). The abnormal threshold was applied using Cytogenomics 2.060 (Agilent Technologies). A combination of several statistical algorithms was applied. A minimum of three oligos that have a minimum absolute ratio value of 0.1 (based on a log(2) ratio) is required for reporting of a copy-number loss or gain. Analysis was performed using Human Genome Build 19 (Feb 2009) as the reference.

## Data access

The sequence and phenotypic data from this study have been deposited in dbGaP (<http://www.ncbi.nlm.nih.gov/dbgap>) under accession number phs000536.v1.p1. The dbGaP accession number assigned to this study is phs000536.v1.p1.

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