A method to sequence and quantify DNA integration for monitoring outcome in gene therapy

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ABSTRACT

Human genetic diseases have been successfully corrected by integration of functional copies of the defective genes into human cells, but in some cases integration of therapeutic vectors has activated proto-oncogenes and contributed to leukemia. For this reason, extensive efforts have focused on analyzing integration site populations from patient samples, but the most commonly used methods for recovering newly integrated DNA suffer from severe recovery biases. Here, we show that a new method based on phage Mu transposition in vitro allows convenient and consistent recovery of integration site sequences in a form that can be analyzed directly using DNA barcoding and pyro-sequencing. The method also allows simple estimation of the relative abundance of gene-modified cells from human gene therapy subjects, which has previously been lacking but is crucial for detecting expansion of cell clones that may be a prelude to adverse events.

INTRODUCTION

Human gene therapy has been carried out successfully for several diseases (1–8), but adverse events have occurred in which subjects developed leukemia associated with insertion of therapeutic vectors near proto-oncogenes (3,6,9,10). For this reason, it is important to track the location and abundance of different integration sites in cells from gene therapy-treated subjects. Tracking integration sites is also of interest in the use of transposons as insertional mutations in model organisms and in basic studies of integrating genomic parasites (11,12).

In the previously used protocols for integration site recovery in gene therapy, gene-corrected cells were isolated from patients, genomic DNA was purified and samples were typically cleaved with restriction enzymes. The exposed DNA ends were then ligated to adaptor DNAs and samples amplified using PCR with one primer complementary to the adaptor and the other complementary to the vector DNA terminus. Sites of integration were identified by sequencing PCR products from a primer bound to the vector DNA, so that the sequence read extended into the flanking human DNA (13,14). It was noticed, however, that integration sites were not equally recovered by different enzymes, pointing to a recovery bias for this method. Although sites were efficiently recovered when found near restriction cleavage sites, they were difficult to detect when not positioned optimally (15,16). Moreover, implementing restriction enzyme-based methods is complex—restriction enzymes need to be identified, which do not cleave DNA within amplicons of interest, only enzymes lacking CpG dinucleotides in their recognition sites can be used due to biased distributions of CpG in mammalian DNA, and
large amounts of genomic DNA are needed. Thus, there is intense interest in alternative approaches.

Improved methods are starting to be described, but all are in early stages (15–18). Concerns with available approaches include efficiency of recovery of rare integration sites (16), and the need for large amounts of genomic DNA for analysis (17,18), which is often not available in gene therapy applications.

Here we report a method for recovering sites of integrated DNA using the bacterial transposase MuA to introduce adaptors into genomic DNA to allow PCR amplification. This method is quick and simple, avoids the bias associated with restriction enzymes, recovers integration sites in a near random fashion, and provides a simple measure of cell clonal abundance.

MATERIALS AND METHODS

Cell lines and gene therapy patient samples

293T and SupT1 cells were infected in vitro using a VSV-G pseudotyped MLV or HIV vector produced by transfecting 293T cells. To generate HIV-based vectors, cells were transfected with the LTR-GFP cassette plasmid p156RRLsin-PPTCMVGFWPRE (19), the packaging construct pCMVdeltaR9 (20) and the vesicular stomatitis virus G-producing plasmid pMD.G. VSV-G pseudotyped MLV particles were produced using pMD.G but in combination with the MLV vector segment (pMX-eGFP) and packaging construct pCGP (pCGP, kindly provided by Paul Bates). Human gene therapy samples consisted of PBMCs (β-thalassemia) or CD3+ cells (SCID-X1). Details of gene therapy samples are reported elsewhere (1,2,7,8). The production of induced pluripotent cells containing a defined number of integration sites is reported in (21).

Mu-mediated integration site recovery

A detailed protocol for preparing Mu reactions is available in the Supplementary Report 1 MuSOP. Briefly, reaction buffer, oligonucleotide donor, target DNA and water are mixed on ice followed by addition of purified Mu transposase. Reactions are then incubated at 30°C for 2–4 h after which 2 μl of the reaction is used as input for PCR. Nested PCR was carried out as described (22) except that extension temperatures during cycling were changed to 70°C for the first seven cycles followed by 67°C for the remaining 37 cycles for samples with HIV-derived vectors. For both HIV- and MLV-based reactions, the number of cycles was reduced to 25 for nested PCR. Single round PCR amplification was performed with primers normally used in the nested round of PCR, each primer encoding 454 adaptor sequences and the LTR primer encoding a DNA barcode (15).

Integration site sequencing and analysis

Integration sites were sequenced using 454 FLX platform technology. Sequences reads were trimmed to remove primer sequence and aligned to the human genome (hg18, version 36.1) using BLAT. Sequences were required to have a single best hit with ≥98% identity to the human genome, to align within 3 bp of the beginning of the sequence read and to contain a perfect match to the expected LTR sequence lying downstream of the barcoded LTR primer. Comparisons to genomic features were carried out as described previously (23,24) using a combination of logistic regression and Bayesian model averaging supplemented by the random Forest machine learning algorithm. Gene expression analyses were based on data from 293T cells (25) with expression measured using the Affymetrix HU133 plus 2.0 gene chip array. Measuring sequence conservation at the site of restriction enzyme or Mu cleavage was performed using weblogo [http://weblogo.berkeley.edu/ and (26)]. Statistical details for generating Lorentz curves and recovery probability plots can be found in Supplementary Report 2 Mu Recovery. Sequence data from this study was submitted to Genbank under accession numbers HR819863–HR863973.

RESULTS

We first quantified biases in the detection of integrated DNA using protocols relying on restriction enzyme cleavage of genomic DNA. Analysis of the recoverability of an integration site based on its proximity to the nearest restriction site showed that integrated DNA ~49 bp from a cleavage site was recovered most frequently, and frequency of recovery decreased sharply at longer or shorter distances (Figure 1A). The case of SCID-X1 patient 10 provides an example of complications due to this recovery bias (3,7). In this patient, an integrated vector activated expression of the nearby BMI1 proto-oncogene, which was associated with massive expansion of leukemic cells. When DNA from blood cells of patient 10 was cleaved and analyzed using four different restriction enzymes, only two enzymes allowed efficient recovery of the BMI1 integration site (Figure 1B). In an extreme effort to circumvent these limitations, a study of SCID-X1 gene-corrected patients used up to six different restriction enzymes to analyze each individual sample, but even with the difficulty and expense of this large scale effort, recovery was still significantly biased (7).

The improved method reported here (Figure 1C and D) substitutes MuA-directed transposition in vitro for restriction enzyme cleavage and adaptor ligation [for background and reviews of MuA function see (11,27–33)]. An engineered transposon end is used as donor (Figure 1C), which contains (i) binding sites for the MuA transposase; (ii) an adaptor region complementary to PCR primers; and (iii) an amine blocking group at one DNA 3’-end. In the presence of MuA transposase, the oligonucleotide donors become covalently joined to target DNA, allowing convenient installation of primer sites in human genomic DNA. PCR amplification is then carried out using primers complementary to the vector DNA end and the adaptor. Because the adaptor contains a 5’ overhang and an aminomodified 3’-end, amplification must begin within the vector DNA and extend through the adaptor, preventing adaptor-to-adaptor amplification. As little as 100 ng of genomic target DNA can be used.
Nested PCR is routinely carried out using the restriction enzyme method, and can be used with the Mu-mediated method. However, tests showed that for samples containing relatively large numbers of integrated vectors, only a single round of PCR was sufficient to yield high quality sequence populations using the Mu-mediated method, and these data sets actually showed improved diversity (described in Supplementary Report 1 ‘Mu Standard Operating Procedure’).

The PCR primers used for the final amplification step are composites, containing both the priming sequences and sequences required for 454/Roche pyrosequencing. Primer sequences also contain a DNA barcode between the 454 sequence and the priming region, allowing large numbers of amplicons to be pooled, sequences determined, then sequence reads parsed using the barcodes (Figure 1D) (34–36). Thus, hundreds of samples can be processed in pools. Pyrosequence reads are then trimmed, aligned to the human genome and distributions analyzed.

To test the Mu-mediated method, we determined 25,194 total integration site sequences, which yielded 3382 unique vector integration sites after condensing duplicates. We analyzed both HIV- and γ-retrovirus-based vectors. Samples studied included cells from patients in two gene therapy trials, which treated SCID-X1 (3) and β-thalassemia (8) and tissue culture cells infected in vitro (summarized in Supplementary Table S1).

We first compared recovery biases of the Mu and restriction enzyme-based methods. We determined the sequence preferences for Mu integration in human DNA in vitro for 5968 integration site sequence reads that included the Mu-end oligonucleotide DNA. Alignment of human sequences at Mu integration sites revealed a detectable consensus sequence closely resembling that reported previously for Mu transposition (37), but with much lower information content than cleavage sites for restriction enzymes (Figure 2A and B and Supplementary Report 2), indicating less bias in the cleaving/joining reactions.

The performance of the restriction enzyme method was next compared with the Mu-mediated method by measuring recovery biases for each method, then annotating the human genome for calculated recovery rates based on the data. Thus, for any integration site in the human genome,
the likelihood of recovery at each base pair could be calculated for each recovery method (Supplementary Report 2).

Figure 3A illustrates the relative recovery frequencies using the LMO2 promoter as an example, which was chosen because the gene has been involved in several adverse events in SCID-X1 gene therapy (3,38). Note that although Figure 3A summarizes results on a single region of the genome, the biases were measured genome-wide from integration site data for each method and are shown at LMO2 for purposes of illustration. Recovery was relatively consistent for the Mu-mediated method over all sites, whereas the restriction enzyme methods show sharp peaks and valleys, where valleys indicate locations where an integration event would be difficult or impossible to isolate and the peaks frequently recovered sites that would mask more rare sites. Figure 3B shows the data plotted as the cumulative recovery frequency, where perfectly unbiased recovery would be indicated by a curve that followed the diagonal from lower left to upper right. The Mu-mediated recovery method most closely approaches the diagonal and is significantly closer than even the method using six restriction enzymes \( (P < 0.001; \text{see Supplementary Report 2 for statistical methods}) \), documenting that the Mu-mediated method is the least biased.

Another means of quantifying recovery biases involves comparing the increased sampling effort required to reach the results of a perfectly unbiased method. The slight biases introduced by the Mu method would require only a 10% increase in sampling effort to achieve the efficiency of a perfectly unbiased method (Supplementary Report 2). In contrast, for some restriction enzyme methods, the needed increase is too large to measure accurately (≥50-fold). For the pool of all six restriction enzymes, a 45% increase in effort would be needed. Thus the Mu-mediated method yields less biased recovery with much less effort than any form of restriction enzyme-based method.

We next investigated the experimental effort required to recover all members of a fully defined integration site population using the Mu-based method. We prepared an induced pluripotent stem (iPS) cell line in which all cells contained the same six known lentiviral vector integration sites. We carried out 12 independent integration site recovery reactions using the Mu-mediated method, recovering an average of 1068 sequence reads per replicate. We found that six of the 12 replicate reactions recovered all six sites. The replicates with fewer than six yielded either four or five of the sites. To assess the sampling effort required for recovery of all six sites, we pooled different numbers of Mu reactions computationally and assessed recovery. For pools of three of the 12 Mu reactions, 98% contained all six sites. For pools of five Mu reactions, 100% of the pools contained all six sites. Thus, 3–5 independent Mu reactions are enough to completely sample an integration site population of this size at the sequencing depth used.

For comparison, three restriction enzymes were tested for recovery of the six sites, and only one yielded all six. The genome-wide distribution patterns of integration target sites for HIV and \( \gamma \)-retroviruses have been studied extensively (13,14,23,39,40), allowing us to assess whether the Mu-mediated method reported similar trends. For studies of integration frequency near genomic landmarks, restriction enzyme-based methods usually provided an adequate overview, because the restriction biases are only weakly related to those landmarks. Figure 4A and B compares the distributions of integration sites isolated using the two methods for HIV and \( \gamma \)-retrovirus-based
vectors. Integration sites are compared from infections of tissue culture cells and for one SCID-X1 patient (3,7).

Figure 4A summarizes the relationship of integration sites to genomic features, using a heat map format to indicate increased or decreased integration frequency compared with random distributions. For both HIV and γ-retroviral vectors, the genome-wide trends were closely similar for the Mu-mediated or restriction enzyme-mediated methods. Both HIV-based and γ-retrovirus-based vectors favor integration in regions of high gene density and associated genomic landmarks. Gamma-retroviruses favor integration near gene 5′-ends. HIV and γ-retroviruses show a complex pattern of favored and disfavored integration sites near regions of histone methylation, acetylation and bound proteins, as indicated by comparison to data from ChIP-seq experiments (41–44) (Figure 4B), again showing favored integration near marks of active transcription. These patterns matched closely for the Mu-mediated and restriction-enzyme-mediated methods. Both HIV-based and γ-retrovirus-based vectors favor integration in regions of high gene density and associated genomic landmarks.

The Mu-mediated method also allows a new approach to quantifying the relative frequency of gene-corrected cells in patient samples. Gene corrected cell clones that are present in many copies will contribute a relatively larger proportion of their integration site DNA to the genomic DNA pool after purification, providing an increased number of target sites for Mu integration in vitro. As a result, relatively larger numbers of independent Mu integration events will lead to recovery of the same high abundance vector integration site. Quantifying the number of independent Mu integration sites per vector integration site therefore provides a simple measure of the relative abundance.

We tested this in a sample from a β-thalassemia gene therapy trial, in which a cell harbouring a single vector integration site in the HMGA2 gene expanded to comprise more than one-third of the gene-corrected peripheral blood mononuclear cells (PBMC) population, as documented by quantitative PCR assays (8). We recovered 10 independent Mu integration sites for the HMGA2 vector integration site, while all other vector integration sites were recovered with only a single Mu site (Figure 4C). This finding makes the important clinical point that the HMGA2 site is the only high abundance integration site in the expanded cell clone. Using the restriction enzyme cleavage method for integration site recovery, only two out of three of the enzymes used allowed isolation of the HMGA2 site (data not shown).

Lastly, we present a method for suppressing PCR contamination. In assessing collections of gene therapy samples, multiple tubes are commonly processed in parallel using nested PCR, providing an ideal setting for migration of PCR products between samples. In practice controlling
contamination is challenging even for experienced experimentalists. We have devised a method that suppresses this, in which separate adaptors are used for each sample, so that contaminating PCR products are not amplifiable outside the correct PCR reaction. In reconstruction experiments, this has been effective at suppressing cross-over in our laboratory (data not shown). Use of multiple adaptors is described in the Standard Operating Procedure in the Supplementary Data for this article.

DISCUSSION

In summary, the Mu-mediated integration site recovery method allows simplified recovery of integration sites and estimation of relative abundance. Use of protocols based on single restriction enzymes results in failure to isolate integration sites that are not near restriction enzyme recognition sites (7,15,16), which can be crucial in monitoring adverse events during gene therapy. The challenges posed by biased isolation have been underestimated in some of the early literature in this field. One previous study attempted to circumvent recovery biases by using six restriction enzymes to study single samples, but even with this added complication and expense, recovery using this method is still more biased than with the Mu-based method (Figure 3B; $P < 0.001$). Thus in cases where it is critical to use integration site data to identify expanded cell clones, the Mu method is attractive and convenient.

Methods for deep sequencing are in a state of rapid transition. The Mu-mediated method described here can in principal be adapted to any of the next generation sequencing platforms. At this writing, the Solexa/Illumina method is least expensive per base, but potentially inconvenient because analysis of a handful of samples will often take up only a fraction of a run, requiring complicated
coordination with others to fill out a run. The 454/Roche method is more expensive per base, but the availability of a benchtop instrument for smaller runs (the ‘Junior’ instrument http://www.gsjunior.com/) simplifies throughput. It is highly likely that additional sequencing methods will become available in the near future that may also be useable with the Mu-mediated method.

The demands placed on integration site recovery technology vary by disease state. The frequency of gene corrected cells varies from 100% (T cells in SCID-X1) to <1% [early adenosine deaminase (ADA) studies], so the demands on the technology differ for different diseases. For very low level clones, recovering integration sites and estimating their abundance is at present challenging for any technology.

Additional methods are starting to be proposed for integration site analysis, including methods based on DNA shearing (N. Gillet, N. Malani, N. Gormley, R. Carter, A. Melamed, D. Bentley, C. Berry, F. Bushman, G. Taylor and C. Bangham, submitted for publication) or limited extension from integrated vectors with a DNA polymerase followed by RNA ligation (‘nrPCR’) (16). Each of these methods is of interest but each may have inefficient steps. For the Mu-mediated method, it can be challenging to obtain enough Mu integration events to query the full human genome efficiently, though the method is suitable for analysis of large numbers of samples with small amounts of starting genomic DNA. Ongoing use has shown the method to be effective in practice. For DNA shearing, it can be challenging to obtain efficient ligation after repairing broken DNA ends, and to work with small amounts of DNA. For the nrPCR method, efficiency may be an issue (16).

Two reports in the peer-reviewed literature document the utility of the Mu-mediated method. In one case, oligoclonal reconstitution during lentiviral vector-mediated gene correction was documented using Mu-mediated recovery of integration sites during β-thalassemia gene correction in mice (46). In this case, the inferred rank order of integration site abundance from sequence read counts was similar to that inferred by quantifying integration site populations.

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**REFERENCES**


**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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