Method to Assemble Genomic DNA Fragments or Genes on Human Artificial Chromosome with Regulated Kinetochore Using a Multi-Integrate System

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Supporting Information

ABSTRACT: The production of cells capable of carrying multiple transgenes to Mb-size genomic loci has multiple applications in biomedicine and biotechnology. In order to achieve this goal, three key steps are required: (i) cloning of large genomic segments; (ii) insertion of multiple DNA blocks at a precise location and (iii) the capability to eliminate the assembled region from cells. In this study, we designed the iterative integration system (IIS) that utilizes recombinases Cre, ΦC31 and ΦBT1, and combined it with a human artificial chromosome (HAC) possessing a regulated kinetochore (alphoidtetO-HAC). We have demonstrated that the IIS-alphoidtetO-HAC system is a valuable genetic tool by reassembling a functional gene from multiple segments on the HAC. IIS-alphoidtetO-HAC has several notable advantages over other artificial chromosome-based systems. This includes the potential to assemble an unlimited number of genomic DNA segments; a DNA assembly process that leaves only a small insertion (<60 bp) scar between adjacent DNA, allowing genes reassembled from segments to be spliced correctly; a marker exchange system that also changes cell color, and counter-selection markers at each DNA insertion step, simplifying selection of correct clones; and presence of an error proofing mechanism to remove cells with misincorporated DNA segments, which improves the integrity of assembly. In addition, the IIS-alphoidtetO-HAC carrying a locus of interest is removable, offering the unique possibility to revert the cell line to its pretransformed state and compare the phenotypes of human cells with and without a functional copy of a gene(s). Thus, IIS-alphoidtetO-HAC allows investigation of complex biomedical pathways, gene(s) regulation, and has the potential to engineer synthetic chromosomes with a predetermined set of genes.

KEYWORDS: human artificial chromosome, HAC, iterative integration system, IIS, DNA assembly, synthetic biology

Since their development, HACs or human artificial chromosomes have been considered to be a promising system for gene delivery and expression, with the potential to overcome many problems caused by the use of virus-based gene transfer systems. Indeed, HACs are stably maintained as single copy episomes. Their use avoids the limited cloning capacity, the lack of copy number control and potential insertional mutagenesis due to integration into host chromosomes that has hampered the use of viral vectors.1−7 There are two types of HACs in general use: “top down” HACs constructed by truncation of the natural chromosomes and “bottom up” HACs generated from BACs carrying 30−200 kb DNA from natural higher-order centromeric DNA repeats. Such BACs are substrates for HAC formation in a process that is accompanied by 20−30-fold multimerization of the input BAC DNA in human cells. Over the past 20 years, many groups have reported the successful generation of HACs of both types1−7 (and references therein), and importantly all HACs constructed so far have constitutive kinetochores.8,9 In the past few years, the amount of research based on the HAC-vectors has significantly increased due to the engineering of HACs with a single loxP gene integration site3,10,11 (and references therein). Such research has included studies on the potential of HACs for gene function analysis, cell reprogramming and animal transgenesis.1−7,12−22

Our group constructed the alphoidtetO-HAC23 from a synthetic alpha-satellite (alphoid) DNA array with precisely defined DNA sequence starting from a 343 bp dimer amplified by the RCA-TAR method23 up to 50 kb in size. This HAC

Special Issue: Genome Engineering

Received: June 14, 2017
Published: August 11, 2017

DOI: 10.1021/acssynbio.7b00209
ACS Synth. Biol. XXX, XXX, XXX−XXX
includes approximately 6000 copies of the 42 bp tetracycline operator (tetO) sequence incorporated into each alphoid dimer. Because the tetO sequence is bound with high affinity and specificity by the Tet repressor (TetR), the 1.1 Mb-size alphoid array in this HAC can be targeted efficiently with TetR fusion proteins that allow the specific manipulation of chromatin within the functional kinetochore. This provides a highly versatile model system for the study of centrochromatin and its impact on kinetochore structure and function in human cells.

In addition, this feature of alphoidtetO-HAC gives the unique possibility to eliminate the HAC from cells, allowing workers to compare the phenotypes of human cells with and without a functional copy of a target gene inserted into the HAC. Such rigorous control is important for proper interpretation of gene complementation analysis and studies of new gene function. Recently the alphoidtetO-HAC was applied to investigate the problem of chromosome instability (CIN), involving the unequal distribution of chromosomes to daughter cells during mitosis which is observed in the majority of solid tumors. While CIN acts as a driver of cancer genome evolution and tumor progression, recent findings point to the existence of a threshold level beyond which CIN becomes a barrier to tumor growth and, therefore, can be exploited therapeutically. Newly developed alphoidtetO-HAC-based assays allow for a quick and efficient screening of hundreds of drugs to identify those affecting chromosome mis-segregation and also to rank compounds with the same or similar mechanism of action based on their effect on the rate of chromosome loss. The fact that the HAC is nonessential in the absence of Blasticidin selection greatly simplifies the interpretation of these assays. Similarly, the alphoidtetO-HAC-based approach was used to identify new human CIN genes, mutations in which are thought to be an early event in tumor development predisposing cells to the accumulation of genetic changes leading to progression to a cancerous state.

A few years ago several laboratories suggested using artificial chromosomes (AC) to assemble entire genomic loci using a multi-integrate system. Such an AC-based approach opens new horizons for synthetic biology, especially in higher eukaryotes. In this study, we designed an iterative integration system (IIS) in such a way that potentially renders it capable of accepting an unlimited number of DNA fragments. In general, IIS is a method to construct large segments of transgenic DNA in a vertebrate host genome. Our IIS method is based on three recombinase enzymes, i.e., Cre, ΦC31 and ΦBT1. Cre is a bidirectional enzyme that catalyzes the recombination between two substrate loxP sites and generates two product loxP sites. In contrast, in the absence of an additional protein Xis, ΦC31 and ΦBT1 are unidirectional enzymes that recombine an attachment bacteria (attB) site and an attachment phage (attP) site to produce attR and attL sites that are not substrates for further reaction. A novel IIS system was combined with alphoidtetO-HAC.

The IIS-alphoidtetO-HAC system can be used to assemble and deliver large DNA fragments that include groups of genes for functional studies. In future, a new HAC-based multi-integrate system carrying multiple genes may be a unique tool for treatment of multigene genetic disorders and for transgenesis experiments with the purpose to understand complex diseases. In addition, the IIS-alphoidtetO-HAC system provides a potentially valuable genetic tool to engineer synthetic chromosomes with a predetermined set of genes in order to investigate complex biomedical pathways and gene(s) regulation.

## RESULTS

### Construction of Plasmids for the IIS-alphoidtetO-HAC System

Five basic plasmids were constructed for the IIS-alphoidtetO-HAC system: the platform cassette A037 on which DNA fragments GENES can be assembled, two carrier vectors used to deliver the transgenic DNA fragments containing promoterless compound markers called either PCF (Pac-mCherry-FcYFur) (Type I A167 plasmid) or GHT (eGFP-hph-TK) (Type II A169 plasmid) and two recombine expression plasmids that express either Cre and ΦBT1 (A135-JH) or Cre and ΦC31 (A139) (Figure 1) (for construction details, see Methods and Figures S1–S5).

The platform cassette A037 consists of the SFM promoter (SV40 enhancer plus Feritin promoter), a split mouse elongation factor 1 (mEF1) intron containing a loxP site within the GHT marker and the attB site of ΦC31 (Figure 1a). The platform cassette was targeted into the unique human Chr13 genomic segment present within alphoidtetO-HAC using homologous recombination in a highly recombinogenic chicken B-lymphoma DT40 cells (Figure 2a). Before targeting, the platform cassette was XhoI-linearized to release 5’ M2A and 3’ M2B hook sequences of 3.5 kb and 3.9 kb in size, respectively, which have homology to the Chr13 genomic segment (see Methods for details). Four pairs of diagnostic primers were used to verify correct insertion of the platform cassette into the Chr13 segment. The expected size products of 4.4 kb and 3.9 kb with B128/B124 and B129/B126 pairs of primers, correspondingly, and 4.6 kb and 4.4 kb with the B034/B126 and B034/B127 pairs of primers, correspondingly, (Table S1) confirms correct recombination between the M2A and M2B hook sequences and homologous sequences of the Chr13 segment (Figure 2b). After insertion of the platform cassette into the HAC, the cells became green (GFP+), resistant to Hygromycin and sensitive to Ganciclovir. Next, the alphoidtetO-HAC carrying the platform cassette was moved from chicken DT40 cells to hamster CHO cells by Microcell-Mediated Chromosome Transfer (MMCT) (Figure 2a). A FISH image of the alphoidtetO-HAC in CHO cells (Clone #CHO BH3:37) is shown in Figure 2c.

The Type I and Type II carrier vectors, A167 and A169, correspondingly, were constructed to deliver a transgenic DNA segment of interest into the platform cassette inserted into the HAC (Figures 1b, 1c). These carrier vectors are YAC-BAC shuttle vectors and, therefore, can propagate in a single molecule state within E. coli as a bacterial artificial chromosome (BAC) with chloramphenicol selection and in S. cerevisiae as a yeast artificial chromosome (YAC) with the yeast HIS3 gene as a selectable marker. Insertion of a transgenic DNA segment into the carrier vector can be done via DNA ligation or yeast-based transformation-associated recombination (TAR) cloning. The Type I carrier vector A167 contains in S'– S' order a loxP site, a promoterless PCF marker, an attB′ ΦBT1 site, a cloning site for DNA insertion, an attP ΦC31 site, a GHT marker under a CAGG promoter flanked by tDNA insulators and a YAC-BAC backbone (Figure 1b). The Type II carrier vector A169 contains a loxP site, a promoterless GHT marker, an attB ΦC31 site, a cloning site for DNA insertion, an attP′ ΦBT1 site, a PCF marker under a CAGG promoter flanked by tDNA insulators and a YAC-BAC backbone (Figure 1c). For the purpose of TAR cloning...
short mammalian genomic DNA segments that do not have yeast ARS-like sequences for a proper propagation in yeast cells, a variant of each carrier vector was made containing yeast origin of replication (ARS), i.e., Type I-ARS (A168) and Type II-ARS (A170) (Figures 1b, 1c and Figures S6, S7). Two expression plasmids were constructed for the IIS-alphoidtetO-HAC system. In these plasmids, a P2A self-cleaving peptide is used to translationally link the expression of two recombinases so that the two recombinases are expressed in equal ratio. The plasmid A135-JH expresses Cre recombinase and ΦBT1 integrase (Figure 1d) while the plasmid A139 expresses Cre recombinase and ΦC31 integrase (Figure 1e). These plasmids also contain a Zeocin marker that is transcriptionally linked to the recombinase expression via an internal ribosomal entry site (ires), allowing selection of these plasmids if desired.

**Description of the IIS-alphoidtetO-HAC System.** The IIS-alphoidtetO-HAC system works as follows. It starts with CHO cells containing alphoidtetO-HAC bearing the platform cassette A037 (Figure 3a). As the GHT marker is expressed, the cells are green (GFP), Hygromycin resistant (hph) and are killed upon exposure to Ganciclovir (TK). Next, these cells are cotransformed with two plasmids, i.e., the A139 plasmid that expresses Cre recombinase and ΦC31 integrase and the Type I carrier vector A167 that contains a transgenic DNA segment of interest (DNA1). Expression of Cre and ΦC31 promotes two recombination events (loxP−loxP and ΦC31attB-attP) between the Type I vector and the platform cassette (Figure 3b). The order of recombination is unimportant as the final product is identical. The SFM promoter within the platform cassette now drives the promoterless PCF marker and contains the adjacent attB′′ ΦBT1 site from the Type I vector, as the original GHT marker is replaced with the PCF marker. This is accompanied by integration of the DNA segment of interest (DNA1) into the platform cassette within the HAC and deletion of all other components of the Type I vector. As a result, the cells that successfully complete both recombination reactions lose green fluorescence and sensitivity to Ganciclovir and gains red fluorescence, resistance to Puromycin and sensitivity to 5-Fluorocytosine.

An error detection mechanism was built into the IIS-alphoidtetO-HAC system because the recombinase-mediated reactions can fail to go to completion. Furthermore, growth, maintenance, screening and storage of candidate vertebrate colonies are far slower, more labor intensive and space limited than either in yeast or bacteria. Thus, it was desirable to remove as many faulty colonies as possible by drug selection and keep only the best candidates for subsequent detailed characterization.

If either of the two recombination reactions fails, this failure event can be selected against and screened out by the error proofing design of the IIS-alphoidtetO-HAC system (Figure 4). As illustrated, the backbone of each carrier vector has its own constitutively active compound marker. Hence, if recombination by Cre fails but ΦC31 occurs (Figure 4a), the A167 Type I carrier vector will integrate into the platform cassette but the PCF marker it carries will remain promoterless. Cells carrying this error are removed by Puromycin selection and counter-selection with Ganciclovir. Alternatively, if recombination by ΦC31 fails but Cre occurs (Figure 4b), the Type I vector A167 will integrate into the construction platform and the SFM promoter will capture the PCF marker, leaving the original GHT marker promoterless. However, the backbone of the

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**Figure 1.** Scheme of plasmids used in the IIS-alphoidtetO-HAC system.
(a) The platform cassette A037. (b) Type I A167 and Type I-ARS A168 carrier plasmids to deliver a genomic DNA fragment. (c) Type II A169 and Type II-ARS A170 carrier plasmids to deliver a genomic fragment. (d) A135-JH plasmid expressing ΦBT1 integrase and Cre recombinase. (e) A139 plasmids expressing ΦC31 integrase and Cre recombinase. Restriction sites used in this work are marked in blue.
Type I vector is retained, and a fully expressed GHT marker under the CAGG promoter remains. Hence, cells generated by such a failure event are Puromycin resistant and have both red and green fluorescence. These types of cells can be removed by counter selection using Ganciclovir against the thymidine kinase component of the GHT marker. In order to avoid the loss of TK gene activity by silencing, the GHT marker is protected by flanking murine tDNA insulators.

Once the first round of recombination is completed (DNA1), the second round of DNA integration (DNA2) can be started. The platform cassette in the HAC now contains a loxP site, an expressed PCF marker and an attB′′ ΦBT1 site (Figure 3c). Expression from the PCF marker gives the cells red fluorescence, resistance to Puromycin and sensitivity to 5-Fluorocytosine. Next, the cells are cotransformed with plasmid A135-JH expressing Cre recombinase and ΦBT1 integrase and the Type II carrier vector A169 containing a second transgenic DNA segment of interest (DNA2). Cre and ΦBT1 expression causes two recombination events (loxP–loxP and ΦBT1 attB′′–attP′′) between the Type II vector and the platform cassette (Figure 3d). This leads to the replacement of the PCF marker by the GHT marker and an attB′′ ΦBT1 site, followed by the insertion of the second DNA segment of interest (DNA2) from the Type1 carrier vector. As a result, the platform cassette in the HAC will contain a loxP site, an expressed GHT marker and an attB′′ ΦBT1 site (Figure 3e). A small insertion (<60 bp) scar comprising the recombination product of attB/attP, the attR site is left between the first (DNA1) and the second (DNA2) DNA segments of interest. Selection with Hygromycin and counter selection with 5-Fluorocytosine ensures that only cells that have correctly undergone the second round of assembly will survive (Figures...
Untransformed parental cells and cells with incomplete recombination are killed by this double selection. After two rounds of recombination, the construction platform is once again where it started, with the exception that two DNA segments (DNA1 and DNA2) of interest have now been integrated into the HAC (Figure 3e). The GHT marker is expressed and the cells have once again green fluorescence, Hygromycin resistance and sensitivity to Ganciclovir. Further rounds of DNA fragment insertions can be repeated indefinitely as required.

Proof of Site-Specific Recombination Using the IIS-alphoidtetO-HAC System. First, we tested whether the ΦBT1 and ΦC31 integrases and Cre recombinase were active and could mediate proper integration into alphoidtetO-HAC carrying the platform plasmid. For this purpose, control experiments were performed using empty vectors. The recombinant assay plasmids were transfected into hamster CHO cells containing the HAC in the combinations A167 plus A139 or A169 plus A135-JH. Each step of insertion was confirmed by PCR.

At the beginning of the experiments, hamster CHO cells carrying the alphoidtetO-HAC bearing the platform cassette were green (GFP+) (Clone #CHO BH3:37). PCR of genomic DNA isolated from the initial CHO cells carrying the HAC gave the expected 778 bp product with the forward primer for the thymidine kinase (TK) gene (B075) and reverse primer for the M2B hook sequence (B681) and no product with the forward primer for the cytidine deaminase gene (FcyFur) (B485) and the reverse primer for the hook M2B sequence (B681) (Figures S8a, S8b; Lane 1) (Table S1).

For the first round of insertion, CHO cells carrying the alphoidtetO-HAC bearing the platform cassette were cotransfected by Type I carrier plasmid A167 and plasmid A139 expressing ΦC31 integrase and Cre recombinase. The cells were cultured in Puromycin/Blasticidin S media. After 10 days of selection, five colonies possessing red but not green fluorescence were picked up and cultured in individual wells in Puromycin/Ganciclovir/Blasticidin S media. In three colonies, insertion of the 2.7 kb Ampicillin resistant gene plus pBR322 DNA fragment from the plasmid A167 into the platform cassette was confirmed by PCR with a pair of corresponding primers: forward B485 and reverse B681 (Table S1). Cells bearing this insertion gave the expected 4.1 kb product with B485/B681 primers but no product with B075/
B681 primers (Figures S8a, 8b; Lanes 2). Clone #E10 was chosen for further experiments.

For the second round of insertion, the cells were cotransfected by Type II carrier plasmid A169 and A135-JH plasmid expressing ΦBT1 integrase and Cre recombinase. The cells were cultured in Hygromycin/Blasticidin S media. After 10 days of selection, ten colonies expressing green but not red fluorescence were picked up and expanded in media containing Hygromycin/5-Fluorocytidine/Blasticidin S. For nine colonies, PCR of genomic DNA isolated from these cells gave the expected 6.2 kb PCR product with B075/B681 primers but no product with B485/B681 primers. This confirmed the correct insertion of the 2.7 kb fragment carried by plasmid A169 (Figures S8a, S8b; Lanes 3). Clone #E10−7 was chosen for further experiments.

For the third round of insertion, the cells were cotransfected and selected for as in the first round. A total number of ten clones were selected. PCR of genomic DNA isolated from these cells gave the expected 6.2 kb PCR product with B075/B681 primers but no product with B485/B681 primers. This confirmed the correct insertion of the 2.7 kb fragment carried by plasmid A169 (Figures S8a, S8b; Lanes 3). Clone #E10−7 was chosen for further experiments.

These results showed that the recombinases are functional and that site-specific insertions into IIS-alphoidtetO-HAC could be achieved.

**VHL Gene Reconstruction Using IIS-alphoidtetO-HAC System.** As a proof of principle, we applied the IIS-alphoidtetO-HAC system to reconstruct the VHL gene. Briefly, the human VHL gene is ~17 kb in size and contains three exons. It is located on chromosome 3 (positions 10137959−10154492; GHCH38/hg38). Mutations in the gene are associated with Von Hippel-Lindau (VHL) syndrome that is a dominantly inherited hereditary cancer syndrome predisposing to a variety of malignant and benign tumors of the eye, brain, spinal cord, kidney, pancreas and adrenal glands.

For the experiments, we used the full-length VHL gene TAR-isolated previously from the total human genomic DNA as a ~25 kb YAC/BAC molecule. Three fragments containing exon 1, exon 2 or exon 3 of the VHL gene were PCR-amplified from the TAR/YAC/BAC clone using specific primers (Table S1). The first Ascl-NotI fragment of 5990 bp in size (positions Chr3:10137959−10143949; GRCH38/hg38) containing exon 1 along with the VHL promoter and the second Ascl-FseI fragment of 6323 bp in size (positions Chr3:10143949−10154492) containing exon 3 were ligated with the Type I carrier vector A168 (Pac-mCherry-FcyFur) that had been digested by Ascl/NotI or Ascl/FseI, correspondingly. The second Ascl-NotI fragment of 4221 bp in size (positions Chr3:10143949−10148169; GRCH38/hg38) containing exon 2 was ligated with the Type II carrier vector A170 (GFP-hph-TK) that was digested by Ascl/NotI (Figure 5a). Three rounds of insertion were performed to assemble the full-length VHL gene using the IIS-alphoidtetO-HAC system.

**Round 1.** A168 vector containing exon 3 and A139 vector expressing ΦC31 integrase and Cre recombinase were cotransfected into hamster CHO cells propagating the alphoidtetO-HAC bearing the platform cassette A037. Cells with this HAC are originally green (GFP+). After the first round, the cells switched to red (mCherry+), Puromycin and 5-Fluorocytidine resistant and carried a modified alphoidtetO-HAC bearing the inserted exon 3 (Figure 5b). A total of 14 clones were obtained after positive selection, with 4 remaining after negative selection: Clone #1, Clone #11, Clone #12 and Clone #14. Clone #14 was chosen for the second round of insertion.

**Round 2.** The A170 vector containing exon 2 and the A135-JH vector expressing ΦBT1 integrase and Cre recombinase were cotransfected into Clone #14e CHO cells carrying the...
VHL gene reconstruction using the IIS-alphoidtetO-HAC system.

**Round 1**
- Construction of Type I A168 and Type II A170 carrier plasmids containing fragments with exon 1 or exon 2 or exon 3 of the VHL gene. Positions of the VHL fragments in the human genome (GRCH38/hg38) are indicated.
- Three rounds of insertion of the VHL fragments into alphoidtetO-HAC carrying the platform cassette are shown. Representative images of changing cell color after each round are shown. (a) Diagnostic PCRs for insertion of the VHL fragments after each round of recombination.

**Round 2**
- Lanes 1 to 5, diagnostic PCR for the fragment 3 with the expected size 1.7 kb. Lane 1, Clone #1; Lane 2, Clone #11; Lane 3, Clone #12; Lane 4, Clone #14; Lane 5, genomic DNA from CHO cells as a negative control.

**Round 3**
- Lanes 6 to 10, diagnostic PCR for junction between fragments 2 and 3. The expected size is 688 bp. The expected size for a negative control is 608 bp. Lane 6, Clone #14−4; Lane 7, Clone #14−12; Lane 8, alphoidtetO-HAC carrying the full-length human VHL gene in CHO cells; Lane 9, a TAR/BAC clone containing the VHL gene; Lane 10, human genomic DNA as a positive control.

RT-PCR for mRNA of VHL. The expected product size is 642 bp. Lane 15, Clone #14−12−3; Lane 16, genomic DNA from CHO cells as a negative control; Lane 17, human genomic DNA from HeLa cell line as a positive control; Lane 18, human genomic DNA from HT1080 cell line as a positive control.
with corresponding primers (first round R3/B678; second round F23/R23; third round F12/R12) (Table S1).

Reconstruction of a functional human VHL gene was confirmed by RT-PCR of the reconstituted RNA transcript from Clone #14–12–3 (Figure Sd) with the primers VHLS-start-F1/VHLEXON-3R (Table S1). The PCR product was sequenced and found to match the correct human VHL sequence (Figure S9). Note that in order for this mRNA reassembly to work, the inserted DNA segments were designed so that the ~60 bp recombination sites were located within introns, and were therefore spliced out of the mature RNA transcript.

**DISCUSSION**

At present, there are many methods to produce transgenic cells for functional studies of genes. The most common methods rely on either transfection of BAC DNA carrying a gene of interest into the host cells or transduction with viruses. However, these methods lead to random integrations into host chromosomes. As a result of that, the expression level of genes varies greatly due to position effects and the number of copies integrated. In addition, the use of viruses limits the size of a gene that can be successfully transduced. Another popular approach is based on integration of a gene into a “hot spot” of a mammalian genome. In this case, homologous recombination for targeting gene integration is very specific due to the usage of a bacteriophage P1-derived Cre recombinase or ΦC31 integrase. However, the efficiency of gene integration remains low. Furthermore, integration of several genes into the same “hot spot” is very difficult, if at all possible.

Several years ago five groups developed a principally new approach to produce transgenic cells.32–36 Their approach was to combine the usage of artificial chromosome-based vectors with a multi-integrase system. This allowed homogeneous gene expression without integration of the vector carrying the target genes into host chromosomes. In addition, artificial chromosomes may be transferred from one cell line to another cell line. Hence, once a chromosome vector expressing a gene of interest was built, it could be moved and used in many different cell lines.

In this study, we developed a novel human artificial chromosome-based system, the IIS-αHAC. This system utilizes two compound markers termed GHT and PCF. Each compound marker is composed of a positive selection marker, a counter selection marker and a fluorescence marker. The GHT marker is composed of sequences encoding green fluorescence protein (GFP), hygromycin phosphotransferase (hph) and herpesvirus thymidine kinase (Tk). The PCF marker is composed of sequences encoding puromycin N-acetyl-transferase (Pac), red fluorescence protein (mCherry), and a fusion protein of cytosine deaminase and uracil phosphoribosyl transferase (FcyFur). These compound markers can be distinguished visually and can be selected for or counter-selected against using the appropriate drug. The IIS-αHAC system includes three enzymes, Cre, ΦC31 and ΦBT1. Binding sites of these enzymes are arranged in a manner that allows the IIS-αHAC to use a promoter capture and a marker exchange strategy to assemble any desired number of genomic DNA segments. In this strategy, the compound marker that is expressed switches each time a new DNA segment is added to the platform cassette of the HAC. The compound markers allow positive selection for cells where a new DNA segment is added to the platform cassette and counter selection against the cells when this event does not happen. Efficiency and accuracy of the IIS-αHAC system has been demonstrated by assembling a functional copy of the VHL gene from multiple DNA segments.

The reassembly of genes from DNA segments by IIS requires breakage and rejoining junctions to be made to the target gene. The optimal placement of these junctions needs to fulfill several requirements. First, the breakage-rejoining junctions must be placed within introns of the target gene. This is necessary as spent integration sites (attR) (<60 bp) are left between adjacent DNA segment (see Figure 3) during the gene assembly process. Placement within introns, allows these integration sites to be spliced out from mRNA and so avoid disruption to the exonic coding sequence.

Second, the placement of breakage-rejoining junctions must not disrupt the splicing efficiency of its host intron. Hence, the breakage and joining junction should not be near any features critical to intron function such as the splice donor site, branchpoint sequence, poly pyrimidine tract and the splice donor site. The spliced donor is on the 5’ end of the intron and this is easily avoided by placing the junction no closer than several hundred bp from it. The branchpoint sequence and poly pyrimidine tract are typically 20–50 bp upstream from the splice acceptor site (3’ end of intron). These sequences can be avoided by placing the breakage-rejoining junction several hundred bp upstream of the splice acceptor sequence. Splice site prediction software can be used to identify the branch sequence (i.e., http://regulatorygenomics.upf.edu/Software/SVM_BP/ or http://www.umd.be/HSF3/).

Third, insertion of the integrase sites at the breakage-rejoining junctions should not generate a cryptic splice donor or acceptor site. Splice prediction software such as http://www.umd.be/HSF3/ can help predict the creation of cryptic splice site.

Fourthly, it is envisioned that TAR cloning would be the preferred method used to clone the gene segments. Hence, the breakage-rejoining junctions must avoid repeat sequences. Repeat sequences can be identified using the “Repeat mask” option in UCSC DNA download window.

Although several laboratories made a significant progress for synthetic biology by constructing multi-integrase systems on different HACs and MACs, a new IIS-αHAC system has some notable advantages that set it apart from other similar artificial chromosome-based systems.32–36 (Figure S10). First, the maximum number of DNA segments that can be added to the IIS-αHAC is only limited by carrying capacity of a human chromosome, which is several hundred Mb. In effect the IIS-αHAC can insert any desired number of genomic DNA segments into the construction platform located within the HAC. Second, each step of insertion is accompanied by a change in cell color that simplifies the selection of correct clones. Third, in this system the insertion “scar” between adjacent DNA segments is greatly reduced (<60 bp), consisting of a single recombinant integration sites (attR) that could, as shown here, be incorporated within introns to allow reassembly a gene from multiple parts. Fourthly, IIS-αHAC has an error proofing mechanism to remove mis-incorporated DNA segments and improve the integrity of assembly. Lastly, as we have previously shown,12,13 an IIS-αHAC carrying a gene(s) of interest can be removed from the cells in culture by targeting with TetR fusion proteins that allow a unique possibility to compare the phenotypes of human cells with and without a functional copy of a gene(s) inserted into the HAC.
For comparison, previously developed artificial chromosome-based systems are limited to the number of efficient recombinases known to work in mammalian cells (see Figure S10a). As a result, the maximum number of fragments that may be inserted into an artificial chromosome is low (five for the systems published and would be approximately 10 if the systems were modified to use all recombinases known to work in mammalian cells). In response, more advanced systems were recently developed by the same group that could perform unlimited multiple insertions (see Figures S10c and S10d). However, all these systems mentioned above retain and integrate the entire plasmid backbone between DNA segments inserted into an artificial chromosome. Therefore, they may experience problems assembling functional genes from gene segments due to the presence of cryptic splice sites or abnormal splicing of the splicing machinery that may change splice isoform generation. Similar problems exist in another artificial chromosome-based system (see Figure S10b). Moreover, as this system uses multiple (<50) identical platform cassettes integrated in the chromosome-vector, the relative position of each new insert relative to one another is uncontrolled. The maximum number of insertions that can be added is limited by the number of platform cassettes, and is further reduced because each round adds an additional loxP site that may recombine and lead to destabilization of the chromosome upon Cre exposure. Another system is based on the principles similar to our system (Figure S10c). However, it has several significant distinctions: it does not include color markers that help to distinguish one round of insertion from another; it does not have an error proofing mechanism; the HAC cannot be specifically and efficiently removed from the host cell cultures.

In summary, the IIS-alphoidtetO-HAC system is able to efficiently and precisely carry out recombination in mammalian cells that allows the investigator to potentially insert any desired number of genomic fragments, leading to assembly of a functional copy of a gene and even more complex loci. The IIS-alphoidtetO-HAC is a valuable unique genetic tool for investigating gene(s) function, complex biochemical pathways and has a great potential for animal transgenesis, development of therapeutic applications for complex diseases, and synthetic biology.

■ METHODS

Cell Lines and Culture. Chicken B-lymphoma DT40 cells containing the alphoidtetO-HAC were cultured in Roswell Park Memorial Institute medium 1640 (RPMI-1640, GIBCO) supplemented with 10% fetal bovine serum (Biowest, Nuaille, France), 1% chicken serum (Invitrogen), 50 μM 2-mercaptoethanol (SIGMA) and 15 μg/mL Blasticidin S (Funakoshi, Tokyo, Japan) at 40 °C in 10% CO₂. Hypoxanthine phosphoribosyltransferase (HPRT)-deficient Chinese hamster ovary (CHO) cells (JCRB0218) carrying the alphoidtetO-HAC were maintained in Ham’s F-12 nutrient mixture (Invitrogen, USA) plus 10% fetal bovine serum (FBS) with 8 μg/mL Blasticidin S Hydrochloride (Funakoshi, Japan). Blasticidin S is required for stable propagation of alphoidtetO-HAC in the medium.

Construction of Plasmids for the IIS-alphoidtetO-HAC System. A037: The platform cassette plasmid was constructed in two parts. The promoter and GHT marker components were amplified from multiple plasmid sources while the recombinate recognition sites, loxP and attB, Flc31 were added by PCR using long oligomers. The GHT compound marker is composed of a fusion of Green Fluorescence Protein (GFP), P2A self-cleaving peptide, Hygromycin phosphotransferase (hph) and viral Thymidine Kinase (TK). The Ch13 targeting hooks (M2A and M2B) were PCR amplified from human genomic DNA and added to the construct. The construction steps and primer sequences used are depicted in Figure S1 and Table S1, respectively.

A167, A168, A169 and A170: The carrier plasmids were constructed in three sections: (i) the integration cassette that comprises of a promoterless compound marker with appropriate recombinase recognition sites; (ii) the error proofing cassette, which is composed of an expressed compound marker opposite to promoterless marker; (iii) the YAC-BAC shuttle vector. Two compound markers were built, the GHT and PCF markers. The GHT marker is as previously described while the PCF marker is a fusion of Puromycin N-acetyl-transferase (Pac), 2A self-cleaving peptide, mCherry fluorescence protein, a second 2A self-cleaving peptide, and cytosine deaminase fused to uracil phosphoribosyl transferase (FcyFur). The construction steps of the Type I carrier plasmid without ARS (A167) and with ARS (A168) are shown in Figure S2 and Figure S6, respectively. The construction steps of the Type II carrier plasmids without ARS (A169) and with ARS (A170) are shown in Figure S3 and Figure S7, respectively. All construction primers used are listed in Table S1.

A135-JH and A139: These are the vectors that express recombinases ΦBT1 and Cre, Flc31 and Cre, respectively. Each vector expresses two recombinases as a single peptide, linked by a 2A-self-cleaving peptide. The construction steps of A135-JH and A139 are shown in Figure S4 and Figure S5, respectively. All construction primers used are listed in Table S1.

Insertion of the Platform Cassette A037 into the AlphoidtetO-HAC in Chicken DT40 Cells. The platform cassette A037 was targeted into the human Ch13 segment present within alphoidtetO-HAC using homologous recombination in a highly recombogenic chicken B-lymphoma DT40 cell line. Before targeting, the platform cassette was digested and linearized by Xhol to release M2A (positions Ch13:69420033–69523541; GRCH38/hg38) and M2B (positions Ch13:69523556–69527457; GRCH38/hg38) hook sequences that have homology to the Ch13 segment. 90 clones were obtained under Hygromycin selection. PCR analysis of genomic DNA from these clones using specific primers (Table S1) confirmed insertion of the platform cassette into the HAC in 14 of the clones.

One Platform Cassette Per AlphoidtetO-HAC, Per Cell. In order for the IIS-HAC system to stably work, only one platform cassette can be present within the HAC and only one cassette within the cell in total. Consequently several strategies and assays were conducted to ensure there was only one platform cassette within the HAC and cell. First, the platform cassette was inserted into a position within the HAC that was previously found to be unique, using homologous recombination in chicken DT40 cell. Targeting was confirmed by PCRs. Hence, there should only be one copy of the platform cassette in the HAC. Second, the HAC containing the platform cassette was then transferred from chicken DT40 to hamster CHO cells by MMCT, thereby removing any copies of the platform cassette that may have integrated into the genome of the DT40 cell. Third, we also conducted fluorescence in situ hybridization (FISH) in conjunction with colony subcloning to ensure that
the CHO cells we worked with had only a single HAC and thus one copy of the platform cassette. Fourthly, the double selection system of IIS innately selects against the presents of more than one platform cassette. Hamster CHO cells, like many cancer cell line, experience chromosome instability. Hence the copy number of the HAC has been observed to spontaneously increase in a small percentage of cells during culturing. This issue is addressed in the IIS-HAC-based system by its compound markers that allow both positive selection and negative counter-selection which are conducted at each round of integration. More specifically, if there were two platform cassettes expressing the GHT marker (GFP-hph-TK) in the HAC, the integration of a DNA fragment with a promoterless PCF (Pac-mCherry-FcyFur) marker into one cassette would result in a cell expressing the PCF marker from one cassette and the GHT marker from the second cassette. The double selection with both puromycin (positive selection for Pac) and ganciclovir (counter-selection against TK) would remove such cells with one or more platform cassettes. Only cells with one platform cassette with the right compound marker can survive rounds of positive selection and negative counter-selection. Lastly and more importantly, PCR analysis of colonies from the integration of “empty vectors” (Figure S8) yielded single bands, indicating the presence of only one platform cassette. If there had been more than one platform cassette within the HAC, after each integration event, we should obtain more than one band in the PCR analysis, i.e., one band for the platform cassette with the insert and one without. Thus, we can conclude that our IIS-HAC-based integration system has only one copy of inserted platform cassette.

**Microcell-Mediated Chromosome Transfer (MMCT) Technique.** The alphoidtetO-HAC containing platform cassette A037 was moved from chicken DT40 cells to hamster CHO cells using an improved microcell-mediated chromosome transfer technique. After MMCT transfer six clones were obtained. PCR of genomic DNA isolated from these clones and FISH analysis confirmed the presence of an autonomous HAC in five clones. Clone #CHO BH3:37 was selected for further experiments.

**Fluorescence In Situ Hybridization (FISH).** FISH analysis was performed as following. Hamster CHO cells carrying the alphoidtetO-HAC bearing the platform cassette A037 were cultured in F12 medium with 10 μg/mL of colcemid (Invitrogen) overnight at 37 °C. Metaphase cells were trypsinized and centrifugated for 4 min at 172g, treated in 10 mL of 50 mM KCl hypotonic solution for 20 min at 37 °C and washed three times in methanol:acetic acid (3:1) solution with a 4 min centrifugation at 172g between each wash. Cells were diluted to the appropriate density with fixative solution, spread onto precleaned slides (Thermo Fisher Scientific, Waltham, MA, USA) above steam (boiling water), and allowed to age 2 days at room temperature. For BAC probing, CHO metaphase slides were washed in 70% formamide in 2x SSC for 2 min at 72 °C. Samples were dehydrated through a 70, 90, and 100% ethanol series for 4 min each and left to air-dry. The probe used for FISH was BAC32–2–mer(tetO) DNA containing 40 kb of alphoid-tetO array cloned into a BAC vector as described previously.11 BAC DNA was labeled using a nick-translation kit with Orange 552 dUTP (S-TAMRA-dUTP) (Abbott Molecular). The probe was denatured in hybridization solution at 78 °C for 10 min and left at 37 °C for 30 min. The hybridization mix probe was applied to the sample and incubated at 37 °C overnight. Slides were washed with 0.4x SSC, 0.3% Tween 20 for 2 min at 72 °C, briefly rinsed with 2x SSC, 0.1% Tween 20 (10 s) and air-dried in darkness. The samples were counterstained with VECTASHIELD mounting medium containing DAPI (Vector Laboratories, Burlingame, CA, USA). Slides were analyzed by fluorescence microscopy. Images were captured using a DeltaVision imaging system in the CRC, LRBG E Fluorescence Imaging Facility (NIH) and analyzed using ImageJ software (NIH).

**Plasmid DNA Transfection and Loading into AlphoidtetO-HAC.** CHO cells with the alphoidtetO-HAC were cotransfected with Type I carrier plasmid A167 and A139 plasmid expressing ΦC31 integrase and Cre recombinase for the first and third rounds of insertion or with Type II carrier plasmid A169 and A135-JH plasmid expressing FBT1 integrase and Cre recombinase for the second round of insertion. Briefly 1 × 10⁶ CHO cells were seeded in one well of a 6-well plate in growth media without selection antibiotics. Next day, 200 μL of Opti-MEM (Gibco), 6 μL of X-tremeGENE 9 DNA Transfection Reagent (Roche) 1.8 μg of A167 and 0.2 μg of A139 or 1.8 μg A169 and 0.2 μg A135-JH plasmids were mixed, incubated 20 min at room temperature and added to the cells dropwise. Next day, the cells were trypsinized and transferred to two 100 mm culture dishes with growth media containing 5 μg/mL Blasticidin S and 5 μg/mL Puromycin in case of the first and third rounds of insertion or 5 μg/mL Blasticidin S and 200 μg/mL Hygromycin B in the case of the second round. The cells were cultured for 7–10 days until colonies of about 1 × 10³ cells formed. Individual colonies were transferred to a 24-well plate and cultured in media with 5 μg/mL Blasticidin S, 5 μg/mL Puromycin and 10 μg/mL Ganciclovir or 5 μg/mL Blastcidin S, 200 μg/mL Hygromycin B and 160 μg/mL 5-Fluorocytidine.

**PCR Reaction.** Genomic DNA was extracted from CHO cultured cells containing the alphoidtetO-HAC with DNeasy Blood & Tissue Kit (Qiagen). DNA concentration was measured by NanoDrop 1000 Spectrophotometer (Thermo Fisher Scientific). The PCR mixture included 0.5 μL TaKaRa LA Taq (5 units/μL), 6 μL of X-tremeGENE 9 DNA Transfection Reagent (Roche) 1.8 μg of A167 and 0.2 μg of A139 or 1.8 μg A169 and 0.2 μg A135-JH plasmids were mixed, incubated 20 min at room temperature and added to the cells dropwise. Next day, the cells were trypsinized and transferred to two 100 mm culture dishes with growth media containing 5 μg/mL Blasticidin S and 5 μg/mL Puromycin in case of the first and third rounds of insertion or 5 μg/mL Blasticidin S and 200 μg/mL Hygromycin B in the case of the second round. The cells were cultured for 7–10 days until colonies of about 1 × 10³ cells formed. Individual colonies were transferred to a 24-well plate and cultured in media with 5 μg/mL Blasticidin S, 5 μg/mL Puromycin and 10 μg/mL Ganciclovir or 5 μg/mL Blastcidin S, 200 μg/mL Hygromycin B and 160 μg/mL 5-Fluorocytidine.
plus 4 \( \mu \text{g/mL} \) Blasticidin S were used as a positive selection for 5 days and 4 \( \mu \text{g/mL} \) Puromycin, 4 \( \mu \text{g/mL} \) Blasticidin S and 5 \( \mu \text{g/mL} \). Gancyclovir were used as a positive/negative selection for 1 week to select for correct insertion of exon 1. All insertions after each round were confirmed by PCR with the corresponding primers (Table S1) that diagnose a correctly assembled VHL gene.

**RT-PCR Reaction.** Transcription of the VHL gene from alpchoi\(^{\text{niu}}\)-HAC/VHL in hamster CHO cells was detected by RT-PCR by using specific primers described in Table S1.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssynbio.7b00209.

Construction history of vectors used in the Iterative Integration System (IIS) (Figures S1–S7); Initial test of IIS and PCR analysis (Figure S8); cDNA sequence comparison between human VHL reconstructed by IIS to native human and native CHO VHL (Figure S9); Schematic diagram of previous integration systems (Figure S10); Primers used in this study (Table S1) (PDF)

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**Author Contributions**

*NCOL and JHK contributed equally to this paper. NCOL initiated and conceived the experiment. NCOL designed and built the integration system. NCOL, JHK, NSP and HSL performed the experiments. NK, HM, WCE and VL analyzed the data. NK and NCOL wrote the paper. WCE and NCO edited the manuscript.*

**Notes**

The authors declare no competing financial interest.

### ACKNOWLEDGMENTS

The authors would like to thank the CRC, LRBGE Fluorescence Imaging Facility (NIH) and personally Drs. Karpova for instructions, consultations and help with the usage of a DeltaVision microscopy imaging system. This work was supported by the Intramural Research Program of the NIH, National Cancer Institute, Center for Cancer Research, USA (VL and NK), a Wellcome Trust Principal Research Fellowship [grant number 107022] (WCE), MEXT KAKENHI grants [16H04747 and 16H01414] (HM) and the Kazusa DNA Research Institute Foundation (HM).

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