Bacterial artificial chromosomes as analytical basis for gene transcriptional machineries

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Abstract Bacterial Artificial Chromosomes (BACs) had been minimal components of various genome-sequencing projects, constituting perfect analytical basis for functional genomics. Here we describe an enhancer screening strategy in which BAC clones that cover any genomic segments of interest are modified to harbor a reporter cassette by transposon tagging, then processed to carry selected combinations of gene regulatory modules by homologous recombination mediated systematic deletions. Such engineered BAC-reporter constructs in bacterial cells are ready for efficient transgenesis in mice to evaluate activities of gene regulatory modules intact or absent in the constructs. By utilizing the strategy, we could speedily identify a critical genomic fragment for spatio-temporally regulated expression of a mouse cadherin gene whose structure is extraordinarily huge and intricate. This BAC-based methodology would hence provide a novel screening platform for gene transcriptional machineries that dynamically fluctuate during development, pathogenesis and/or evolution.

Keywords Bacterial artificial chromosome (BAC) · Transgenic mouse · Gene transcriptional regulation · Enhancer · Transposon · Homologous recombination · Cdh6 (cadherin-6)

Introduction

In the post genome-sequencing era, genome-wide means of functional assay has become possible to systematically understand principles of the life and its changeful conditions (i.e., homeostatic regulations, disorders etc.). For example, by taking advantage of the DNA microarray screening system, one can describe which genes among tens of thousands are transcribed into mRNAs at any time and place of the developing and/or pathogenic organisms (reviewed in Dufva 2009; Lockhart and Winzeler 2000). The proteomics standard by which extremely sensitive detection capacity of mass spectrometry can characterize every small protein spot from multiple dimensional electrophoresis has also opened a door for high-throughput means of profiling translated protein dynamics in various contexts of organism’s development and/or life conditions (Choudhary and Mann 2010; Pandey and Mann 2000 and references therein). Regarding the gene regulatory machineries, bioinformatics tools such as evolutionarily conserved region (ECR) browser (Ovcharenko et al. 2004) help list the non-coding genomic fragments highly conserved among species, which might be implicated to
play significant roles during organism’s development, pathogenesis and/or evolution. The chromatin immunoprecipitation (ChIP) based technology (i.e., ChIP-sequencing; Robertson et al. 2007) further allows identification of those genomic fragments with which given transcriptional protein complexes interact and the ENCODE project (Thomas et al. 2006) recently unifies such comparative/experimental information on the whole human genome, providing a useful starting point to examine gene regulatory landscapes (reviewed in Natoli 2010). However, gene transcriptional regulations are elaborated by synergistic interactions among these functional genomic fragments that are sometimes scattered over a mega-base sized genomic territory and it is still of challenge to quickly and thoroughly investigate the value of non-coding genomic regions from any species in any in vivo contexts.

Bacterial Artificial Chromosome (BAC) contains an Escherichia coli derived F-factor replication origin, allowing stable propagation of a large exogenous DNA fragment (average size: ~200 kb) by a single copy per a bacterial cell in a supercoiled circular form (Shizuya et al. 1992; Shizuya and Kouros-Mehr 2001). Yeast Artificial Chromosomes (YACs) could also keep mega-base sized DNA fragments in a long linear structure (Burke et al. 1987), yet YAC inserts are often chimeric with a higher rate of recombination, making it difficult to steadily engineer and maintain the YAC structure all the time (Shizuya et al. 1992; Shizuya and Kouros-Mehr 2001). From this reason, BACs rather than YACs had been utilized as rigid minimal components of genome-sequencing projects, and researchers can now easily obtain BAC clones that differentially cover any genomic regions of human, mouse, rat, chicken and so on. Recently, resourceful methods to precisely manipulate BAC clones by means of simple homologous recombination as well as transposon tagging in bacterial cells were developed (Yang et al. 1997; reviewed in Copeland et al. 2001). Since BAC transgenic system is efficient in wide variety of cell lines and fertilized eggs from mouse, frog and zebrafish (Poser et al. 2008; Montigny et al. 2003; Wade-Martins et al. 2001; Antoch et al. 1997; Kelly et al. 2005; Jessen et al. 1998), setting BACs as analytical basis would be a promising way for functional genomics at the post genome-sequencing era. Many research groups including ours (Spitz et al. 2003; Jeong et al. 2006; Inoue et al. 2008; Carvajal et al. 2008) have certainly engineered BACs to survey large cis-regulatory landscapes in mouse. Nonetheless, investigative principles and/or conditions for BAC modification toolkits are even now mixtures of good and bad and extensive adjustments would be required for their effective use.

In the present study, we introduce an experimental strategy in which mouse or virtually any vertebrates’ gene regulatory regions can be characterized from a mega-base sized genomic territory more comprehensively than ever. As the first step, transposon mediated reporter tagging was processed among BAC clones that differentially cover the genomic territory of interest in bacterial cells. These BACs are then microinjected into fertilized mouse eggs to select BAC clones that can recapitulate gene regulatory patterns of interest. Next, homologous recombination mediated systematic regional deletions from above selected BAC-reporter construct are carried out in bacterial cells and these modified constructs are again microinjected into mouse fertilized eggs to decisively evaluate the reporter activity in the developing mouse organisms: in this final step, rigorous assessment of how BAC deletion patterns affect reporter expression profiles reliably distinguishes critical regions for gene transcriptional regulations. By applying these methodological combinations, we could quickly identify a gene regulatory region for spatio-temporally restricted expression of a mouse cadherin gene whose organization had been too huge and complex to examine gene regulatory machinery with conventional methodology and/or bioinformatics. This strategy would thus provide a novel high-throughput screening platform for any non-coding genomic fragments that might harbor unknown significance in multi-cellular organisms’ development, evolution and/or pathogenesis.

Materials and methods

Transposon mediated reporter tagging of BAC clones

A modified version of Tn1000 carrying the BGZ40 cassette used in this study was described previously (Morgan et al. 1996). Briefly, BAC clones RP24-88H4, RP23-190L12 or RP23-78N21 that differentially cover mouse cadherin-6 (Cdh6) gene locus

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were electrooporated into the male strain *MH1844* in which modified *Tn1000* is resident on the sex factor plasmid *R388* (Morgan et al. 1996). Next, the BAC transformant was mated with the Streptomycin (Strp) resistant female strain DH10B on LB agar plate for 2 h at 37°C and those females received modified BAC with single random transposon insertion were selected on LB agar plate containing 100 µg/ml Strp and 12.5 µg/ml Chloramphenicol (Cam) at 37°C. After getting rid of *R388* from selected colonies by re-electroporation into DH10B strain, positions of transposon insertion were determined by direct sequencing of modified BACs from both ends of the transposon (Primers for direct sequencing were described in Morgan et al. 1996).

Homologous recombination mediated modification of BAC clones

A recombinogenic bacterial strain EL250 was used as previously described (Lee et al. 2001). For homologous recombination mediated BAC deletions, ~1-kilo-base pair (kb) sized 5′ and 3′ arms were amplified via polymerase chain reaction (PCR) and cloned into the *pGEM-T-easy* vector (Promega) and in between these arms, an Ampicillin (Amp)-resistant (r) gene cassette or a Kanamycin (Kan)-r cassette put between two FRT sites was subcloned. Then the portion of arms with a selection cassette was released from the vector and kept at −30°C. Homologous recombination for BAC deletion was processed as was shown in Figs. 3a and 4b. Briefly, BAC clones for modification were electrooporated into the recombinogenic EL250 strain and selected on L-agar plate containing 12.5 µg/ml Cam. Next, a selected single colony was inoculated in 50 ml of LB without antibiotics at 32°C until the OD 600 reached to 0.5 and 15 ml of the culture was heat-shocked at 42°C for 15 min, while another 15 ml was maintained at 32°C for 15 min as negative control. Both cultures were placed on ice for 15 min, centrifuged and quickly washed by ice-cold water to keep the electro-competency and recombination activity. A few hundred nanograms of the arm-selection cassette fragment was then electrooporated into the electro-competent cells and positive selection was carried out at 32°C on L-agar plate containing 12.5 µg/ml Cam and 12.5 µg/ml Kan or Amp for ~40 h. Ordinarily, as many as 100 colonies made growth from the heat-shocked competent cells, while few colony growth occurred from the control ones. After the homologous recombination, Amp-r or Kan-r cassette was excised by adding arabinose to the log phase liquid culture of the EL250 cells with the correctly modified BAC at 0.1% v/v for 1 h at 32°C to sufficiently induce Flpe (Fig. 3a). For confirmation of the deletion and excision, pulse filed gel electrophoresis (PFGE) by using the CHEF-DRII system (BioRad) and PCR based evaluation were performed. As for PFGE, NotI digested BAC constructs were applied in 1% agarose gel containing 0.5 x Tris-borate-EDTA (TBE) buffer for electrophoresis and the gel was placed under 6 V/cm electric field in cooled 0.5 x TBE buffer circulation with the field switch time being set at 1–20 s for 19 h. Small blocks of MidRange 1 PFG Marker (#N3551S, New England Biolabs) were simultaneously applied to the gel for estimation of the BAC fragment sizes.

Generation of BAC transgenic mice

All animal experiments in this study conform to Japanese governmental guidelines and have been approved by the Animal Care and Use Committee of the National Institute of Neuroscience, Japan (Project #2007022). BAC transgenic mice were generated and genotyped as previously described (Inoue et al. 2008). Briefly, BAC DNA was purified from 1.5-1 LB bacterial culture by equilibrium ultracentrifugation in continuous CsCl-ethidium bromide gradients. The purified BAC was linearized by PI-SceI and dialyzed to the injection buffer. Final concentration and purity of the linearized BAC-reporter construct was checked by an O.D. meter (NanoDrop 1000, Thermo). For microinjections into pronuclei of fertilized eggs derived from the superovulated B6C3F1 mouse strain (Charles River, Japan), DNA was diluted to ~ 2 ng/µl by the injection buffer and tails of transgenic founder (F0) pups or yolk sacs of F0 embryos were collected for the genotyping. For BAC transgenic founders/embryos, presence of *RP23/24*-BAC vector sequences immediately upstream and downstream of the PI-SceI site was always confirmed by PCR to minimize the possibility that fortuitous deletions fell on the BAC construct integrated into the mouse chromosome as was described in Inoue et al. (2008).
Whole-mount detection of beta-galactosidase activities in the transgenic embryos

Timed pregnant mouse embryos precisely staging by Theiler (1972) were stained as described in Inoue et al. (2008). Photographs were taken under the binocular microscope (MZ8, Leica) equipped with a CCD camera (DFC300 FX, Leica).

Results and discussion

Transposon mediated reporter cassette insertions into a given BAC clone effectively trapped gene transcriptional machineries included within the BAC

Transposon is a powerful DNA delivery tool if its insertion and excision can strictly be controlled. In order to trap gene regulatory machineries included within a single BAC clone, we utilized a modified version of the Tr1000 bacterial transposable element in which a copy of BGZ40 reporter cassette was stably introduced (Fig. 1a; Morgan et al. 1996). The BGZ40 cassette contains a human beta-globin minimal promoter, Escherichia coli beta-galactosidase gene (LacZ) as a reporter, and a polyadenylation signal (pA) from simian virus 40 (SV40), virtually allowing rigid reporter activation after the transgenesis only when gene regulatory elements (i.e., enhancers) interactive with the human beta-globin minimal promoter were included in the modified BAC clone.

We initially examined how the BGZ40 modified transposon was integrated into a single BAC clone RP24-88H4 that mainly covers 5' upstream region of the mouse cadherin-6 (Cdh6) gene transcription start site (TSS; Fig. 1b). As was previously reported, transposon mediated integration of the reporter cassette into BACs can be achieved via simple 2 h’ interstrain mating of bacterial cells (Morgan et al. 1996) and we indeed found that every selected colony after the mating process harbors a single insertion of transposon at quasi random positions in the BAC clone (Fig. 1b). We usually obtained hundreds of colonies from the bacterial mating mix derived from a single 100-mm plate dish with ~90% of the colonies yielding right transposon insertions. This demonstrates that a single mating experiment for a single BAC clone is good enough to collect a series of reporter modified BAC clones by the transposon tool.

We then selected four reporter modified RP24-88H4-BAC clones (#166, #207, #208, #209; Fig. 1b) and generated transgenic mouse lines from each BAC to evaluate the reporter activity in vivo. As the results, we confirmed that all these four clones recapitulated exactly the same reporter expression patterns along differentiating Schwann cells (white arrows in Fig. 1c–f; Table 1), some other neural crest derivatives homed in branchial arches (white asterisks in Fig. 1c–f) as well as at the trunk region (Fig. 1c–f), mesenchymal cells in the limb buds (Fig. 1c–f) and cells along the dorsal midline at embryonic day e12.5 (Fig. 1c–f). This indicates that the BGZ40 reporter cassette is capable to capture some of the activities of gene regulatory elements included in the BAC clone RP24-88H4 regardless of its integration sites.

We further evaluated to what extent the BGZ40 cassette can detect the activity of gene regulatory elements included in a given BAC clone. For this purpose, we chose another BAC clone RP23-78N21 that covers both the Cdh6 gene TSS-exon and ATG-exon to carry out two types of modifications on this clone: One is the transposon mediated insertion of BGZ40 cassette into the BAC clone and the other is the recombination mediated in-frame insertion of the LacZ-SV40pA cassette at the Cdh6-ATG-exon which is included in the BAC clone. In the latter configuration, the native Cdh6 promoter element located at around the TSS is expected to activate the reporter gene in the modified BAC clone, logically capturing activities of all gene regulatory elements included in the clone after the transgenesis (Fig. 2a). As was shown in Fig. 2a, we prepared one transposon modified BAC clone and one homologous recombination modified BAC clone to generate transgenic mouse lines from each BAC-reporter construct. As the consequence, we found that e12.5 transgenic embryos from these two modified BACs recapitulated very similar reporter expression patterns along the neural crest derivatives at the trunk region as well as in the brainchial arches and so on (Fig. 2b, c; Table 1). Noticeably, expression levels of LacZ from the homologous recombination modified construct tended to be higher than those from the transposon modified construct probably due to the promoter interference
between endogenous Cdh6 promoter and BG minimal promoter and/or the intron-less feature of the BGZ40 cassette (Fig. 2b, c). The result demonstrates that the human beta globin minimal promoter in the BGZ40 cassette is good enough to detect virtually all activity of gene regulatory elements included in a given BAC clone after the transgenesis in mice.
Homologous recombination mediated systematic deletions from the transposon-modified BAC clones pinpoint the gene regulatory modules

We next sought to establish a reliable assay system in which systematic deletions of given genomic fragments from the reporter modified BAC clone by means of homologous recombination in bacterial cells. Since the transposon modified BAC RP24-88H4 clones fairly recapitulated the Schwann cell specific expression of Cdh6 gene at e12.5 (Fig. 1c–f), we tried to confine which genomic fragments in the BAC clone is required for the restricted gene expression. In the previous report, we selected five BAC clones that differentially cover the ~350-kb genomic territory of huge Cdh6 gene locus and found that each BAC with reporter modification fairly recapitulated distinct combinations of Cdh6 mRNA expression profiles after the transgenesis (Inoue et al. 2008). From these results, we had already speculated that ~36-kb genomic fragment covered with BAC RP24-88H4 might be critical for Cdh6 expression in Schwann cells (Inoue et al. 2008). We thus designed to generate a BAC based reporter construct in which ~36-kb genomic fragment is completely deleted from a modified RP24-88H4 clone #166 and examined if ~36-kb genomic region is required for the reporter expression in Schwann cell populations after the transgenesis. For the deletion, we subcloned 1-kb 5’ upstream and 1-kb 3’ downstream fragments from the target region as homologous arms for recombination and further placed an Amp-r cassette in between the arms. The configuration of this construct logically allows precise excision of the ~36-kb genomic fragment from the BAC clone after the heat shock inducible homologous recombination event in the recombinogenic Escherichia coli strain EL250 (Fig. 3a, b). As the results, we obtained the right BAC-reporter construct with high efficiency (i.e., 4 colonies out of randomly selected 6 colonies contained the correctly modified clone), which was verified by PCR and pulse field gel electrophoresis (PFGE; Construct #2 in Fig. 3b, c).

To further make sure if any types of homologous recombination mediated BAC deletion can be processed with reasonable efficiency, we tried to generate several series of deletion constructs in which various size and/or combinations of deletions are introduced into the reporter modified BAC clones (Fig. 3b). Consequently, we confirmed by means of PCR and PFGE that any types of deletions can equally be placed into a given BAC clone (Fig. 3c, d). For instance, recombination efficiency for ~400-base pairs (bp) and 77-kb deletions appeared to fall at a similar level (i.e., 4–5 colonies out of randomly selected 6 colonies contained the correctly modified clone), which was verified by PCR and pulse field gel electrophoresis (PFGE; Construct #2 in Fig. 3b, c).

Table 1 Summary of Tg mouse lines analysed in the present study

| Figure | BAC transgene | Independent Tg mouse lines generated | Tg with the reproducible expression pattern as in Figures | Tg with the expected expression pattern |
|--------|---------------|-------------------------------------|--------------------------------------------------------|----------------------------------------|
| 1-C    | RB24-88H4, Tn:BGZ40 #207 | 9 | 7 | 7* |
| 1-D    | RB24-88H4, Tn:BGZ40 #208 | 8 | 6 | 6* |
| 1-E    | RB24-88H4, Tn:BGZ40 #209 | 9 | 8 | 8* |
| 1-F    | RB24-88H4, Tn:BGZ40 #166 (Fig. 3b #1) | 7 | 6 | 6* |
| 2-B    | RP23-78N21, Rec:LacZpA | 7 | 6 | 6 |
| 2-C    | RP23-78N21, Tn:BGZ40 | 2 | 2 | 2 |
| 3-B    | RB24-88H4, Tn:BGZ40 #166 Δ36 kb (#2) | 7 | 7 | 0* |
| 3-B, 3-E | RB24-88H4, Tn:BGZ40 #166 Δ10 kb (#3) | 3 | 2 | 2* |
| 3-B    | RB24-88H4, Tn:BGZ40 #166 Δ10 kb (#4) | 7 | 7 | 0* |
| 3-B, 3-F | RB24-88H4, Tn:BGZ40 #166 Δ400 bp (#5) | 4 | 4 | 0* |

* Expression in Schwann cells, Tg Transgenic
We finally purified and transferred the BAC construct #166 with ~36-kb deletion (=Construct #2 in Fig. 3b) into mouse fertilized eggs to obtain transgenic mouse lines. As the results, we found that the reporter expression in Schwann cell populations entirely disappeared from these mouse lines, while the expression in other regions such as neural crest derivatives in the trunk and cranial branchial arch was unchanged at e12.5 (Table 1; T. Inoue et al., manuscript in preparation). This indicates that the 36-kb territory is indeed required for the Cdh6 expression in Schwann cell populations at e12.5. Other deletion BAC constructs except for construct #3 also abolished reporter expression in Schwann cell lineage after the transgenesis, further demonstrating that a ~400-bp module in the 36-kb territory that contains an ECR with several transcription factor binding sites is critical for the Cdh6 expression at e12.5 (Fig. 3e, f; Table 1; T. Inoue et al., manuscript in preparation). Since the sets of BAC deletion constructs described in this study were able to be simultaneously prepared for the transgenesis within 3–4 weeks by a researcher, our strategy might provide a high-throughput screening platform for gene transcriptional machineries that dynamically fluctuate during development, pathogenesis and/or evolution.

A comprehensive strategy in characterizing gene transcriptional machineries

In the conventional method for examining gene transcriptional machineries (additive strategy: Fig. 4a), genomic fragments whose size is limited to several kilo-base pairs in plasmid constructions are linked to the reporter cassette and the reporter activities are monitored in cell lines and/or transgenic animals. In order to analyze a genomic region whose size is longer than several hundreds kilo-base pairs, one should hence prepare more than hundreds of plasmid based constructs to identify sufficient modules for spatio-temporally restricted gene expressions (Fig. 4a). While recent bioinformatics based predictions greatly help confine possible transcription factor binding modules in the genome, it is still needed to prepare tens of plasmids in verifying the physiological significance. Moreover, this strategy cannot identify those gene regulatory modules that are separated each other into a large genomic region yet synergistically work together for the gene activation/silencing.

From our experimental results described above, we now propose a systematic way (subtractive strategy: Fig. 4b) for narrowing down transcriptional regulatory
Fig. 3 BAC-reporter construction with efficient homologous recombination mediated deletions. 

**a** The flow chart to introduce deletions into BAC-reporter construct by homologous recombination. Cam Chloramphenicol, Kan Kanamycin, Flpe Flipase, FRT Flpe recombination target, r resistant gene.

**b** Examples of BAC-reporter deletion constructs generated by homologous recombination. Constructs #2–#5 were generated from the transposon tagged RP24-88H4 (Construct #1 is identical to Construct #166 in Fig. 1b), while Constructs #7–#9 were originated from the transposon tagged RP23-190L2 (Construct #6). The size of deletion is depicted with the selection cassette used for the homologous recombination. Amp-r, ampicillin resistant gene cassette; Tn:BGZ40, integrated LacZ reporter cassette via transposition.

**c** Band patterns confirmed by pulse field gel (PFG) electrophoresis. Each BAC-reporter construct was digested with NotI and the fragment size was estimated by comparison with the PFG markers (M) whose size components were indicated between the panels. **Bold characters** correspond to the **bold marker bands**.

In panel **c**, a band in lane #1 referred by red asterisk is modified by Amp-r integration. In panel **d**, a band in lane #6 referred by red asterisk is modified by Amp-r integrations, while that in lane #6 referred by green asterisk is modified by Kan-r integrations. Note that bands referred by corresponding asterisks in lanes #2–5 and #7–9 correctly shift with BAC modification patterns summarized in panel **b**. **e**, **f** Representative reporter expression patterns after the transgenesis of Construct #3 (e) and #5 (f). Note that the reporter expression in differentiating Schwann cells (white arrows) was totally abolished in BAC transgenic mouse embryos generated from Construct #5 (f), indicating value of the deleted 400-base pair (bp) sized region for the expression. **White asterisks**, expression in the branchial arch.

The ratio of embryos exhibiting reproducible reporter expression over the total number of independent transgenic mouse lines generated is noted at the left bottom corner of each panel (e, f) and in Table 1.
machineries into short territories from large genomic sections. As the first step, transposon mediated reporter integration into BAC contig is to be set as the analytical basis for enhancer screening. For instance, given that a ~0.5-mega-base pair (Mb) sized genomic territory arises as the screening target for a gene regulatory machinery of interest, four BAC clones would be enough to differentially cover the territory (Constructs a–d in Fig. 4b) and it is straightforward to modify all of them by transposon tools compared to the recombination-based method that requires inclusion of transcription start sites in the target BAC clones. This could primarily capture almost all enhancer activity in these BAC clones (refer to Fig. 2) and if one can find out some of them recapitulated the gene expression patterns of interest after transgenesis, essential regulatory modules are speculated to be distributed within a ~50-kb genomic segment (Fig. 4b; In this example, one can speculate that segment III is required for the reporter expression because only Construct b yields the expression after the transgenesis). As the second step of evaluation, recombination mediated systematic deletions from the reporter modified BAC clone would be the most comprehensive way in identifying a critical genomic fragment from the speculated ~50-kb genomic segment for the transcriptional regulation of interest. For instance, if two recombination mediated deletion constructs were generated from the reporter modified BAC clone with the deleted region being equally overlapped each other (Construct b’ and b” in Fig. 4b), one can narrow the ~50-kb segment into a fragment whose size is as short as the one third of the segment by examining the reporter activity in the transgenic animals (Fig. 4b; In this example, territory A in the segment III is speculated to be essential for the reporter expression because only Construct b” yields the expression after the transgenesis). It should be of note for this step that introduction of ‘two’ equally
overlapped deletions to a target segment of a selected BAC-reporter construct is a key to minimize experimental efforts: two equally overlapped deletions could narrow the critical region for transcriptional regulations into 1/3rd of the target segment (Fig. 4b), and two more cycles of experiments with similar deletion approaches would further confine the region into \( (1/3)^3 = 1/27 \) th of the target segment. On the other hand, three equally overlapped deletions could confine the regulatory regions into 1/5th of the target segment and one more cycle of experiment would restrict the regions into \( (1/5)^2 = 1/25 \) th of the target segment. Considering the facts that both examples described above require six BAC-reporter deletion constructs for the screening and that more overlapped deletions need more constructions, two-overlapped-deletion based screen is certainly the most efficient way to finally pinpoint the gene regulatory modules.

Taken together, only six BAC-reporter constructs (Constructs a–d, b’ and b” in Fig. 4b) and their transgenesis stand logically sufficient to narrow the 0.5-Mb genomic territory into ~15-kb genomic fragment for gene transcriptional regulation (Fig. 4b) and here proposed new strategy indeed allowed us to identify many of Cdh6 gene regulatory modules which cannot easily be examined by the conventional method for its huge and complex genomic organization (Fig. 3; our unpublished results).

Generally, the subtractive way of screening in characterizing critical factor(s) from tens of pooled candidates is a clear-cut and efficient option. For the immediate example, the four Yamanaka-factors that cooperatively induce pluripotent stem cells from somatic fibroblasts have been comprehensively identified by simultaneous gene transfections of any 39 factors among the pooled 40 candidate factors (Takahashi and Yamanaka 2006). In this case, only 40 rounds of experiments with ‘subtractive’ pools quickly solved out the four indispensable factors from possible trillions of combinations (=240). Hundreds of candidate gene regulatory modules scattered over a given genomic territory can be considered as the analogous situation. Notably, BAC clones derived from human or chick genomic library can similarly be modified and evaluated in mouse system after the transgenesis (our unpublished observation). Spitz et al. (2003) and Jeong et al. (2006) have already reported similar screen strategies for gene regulatory regions, yet the present study further confirms the utility of BAC modification tools in evaluating gene regulatory elements (Figs. 1, 2, 3) and improves the way of evaluation that never primarily depends on weighting with bioinformatics and/or earlier experimental evidence-biased predictions (Fig. 4). Kokubu et al. (2009) have also innovated the transposon-based mouse chromosomal engineering method to broadly map cis-regulatory landscapes for any gene clusters. While this strategy surpasses transgenic-based systems including ours in the point that it allows integration of the reporter cassette into the natural genomic context, it would be time/money-consuming to attain the kilo-base-pair-sized analytical accuracy only with their local-hopping-transposon-tagging and deletion strategy in mouse embryonic stem cells. Our BAC-transgenic based methodology with systematic deletions (=subtraction of gene regulatory modules) in bacterial cells might complement the inventive Kokubu’s method and would thus provide ideal foundations in assessment of any detailed gene regulatory landscapes most of which remain elusive.

Conclusions

The great majority of vertebrate genome is occupied by the non-coding DNA sequences that are thought to be involved in regulation of chromosomal structures and/or differential gene expressions. Mobile DNA elements and some other repetitive DNA sequences resident in the non-coding regions are also known to affect gene expression profiles. Recent studies have further revealed that human copy number variations as well as single nucleotide polymorphisms in the non-coding regions tightly link to pathogenic conditions. It must therefore be essential to be able to systematically annotate these non-coding DNA sequences and our BAC based methodology should help establish a novel high-throughput screening platform for these sequences that might harbor unspecified yet critical roles for organisms’ development, pathogenesis and/or evolution.

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