

Big DNA as a tool to dissect an age-related macular degeneration haplotype

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Abstract

Age-related Macular Degeneration (AMD) is a leading cause of blindness in the developed world, especially in aging populations, and is therefore an important target for new therapeutic development. Recently, there have been several studies demonstrating strong associations between AMD and sites of heritable genetic variation at multiple loci, including a highly significant association at 10q26. The 10q26 risk region contains two genes, *HTRA1* and *ARMS2*, both of which have been separately implicated as causative for the disease, as well as dozens of sites of non-coding variation. To date, no studies have successfully pinpointed which of these variant sites are functional in AMD, nor definitively identified which genes in the region are targets of such regulatory variation. In order to efficiently decipher which sites are functional in AMD phenotypes, we describe a general framework for combinatorial assembly and delivery of large 'synthetic haplotypes' to relevant disease cell types for downstream functional analysis. We demonstrate the successful and highly efficient assembly of a 119kb wild-type 'assemblon' covering the *HTRA1/ARMS2* risk region. We further propose the parallelized assembly of a library of combinatorial variant assemblons covering the region, delivery and analysis of which will identify functional sites and their effects, leading to an improved understanding of AMD development. We anticipate that the methodology proposed here is highly generalizable towards the difficult problem of identifying truly functional variation discovered via GWAS or other genetic association studies.

Introduction

Age-related macular degeneration is a complex genetic disease characterized by common haplotype associations.

Age-related macular degeneration (AMD) is considered to be the leading cause of vision loss in the elderly of developed countries (1-3). As the populations of these countries continue to age, AMD will increase in prevalence, and its already significant contribution to health care costs will continue to grow. The disease is characterized by abnormalities in the retinal pigmented epithelium (RPE), which underlies the photoreceptor cells of the retina (**Fig. 1**) (4). AMD primarily affects the macula, a region of the central retina associated with the highest acuity in the visual field. While the etiology of AMD is complex and unclear, the disease has a well described progression from early, to late, and on to advanced forms. The advanced forms of the disease are the primary cause of vision impairment due to AMD, with 90% of patients exhibiting severe central vision loss having progressed to advanced stages of disease (5).

Despite the well-known progression of disease stages, AMD is not curable, and current therapeutic interventions have had limited effectiveness, owing largely to its complex genetics and incomplete knowledge of the disease's underlying mechanisms. A combination of high-dose zinc along with vitamins C and E and β -carotene is a common treatment recommended for newly diagnosed patients (6). The treatment has no effect on developing AMD but may have some benefit for slowing progression of AMD once it has started. Vascular endothelial growth factor (VEGF) is a major factor associated with new blood vessel growth and is thus expressed in the neovascular membranes of wet AMD. Anti-VEGF therapies have therefore been used as some of the most effective treatments for CNV, with up to 90% of patients seeing stabilized vision and nearly a third reporting significantly improved vision in some clinical trials (7, 8).

Both environmental and genetic risk factors have been implicated in the development and progression of AMD. Smoking is the strongest modifiable environmental contributor to AMD, and has been shown to increase the overall risk of AMD diagnosis by two- to three- times (9). Environmental risk factors are likely only modifiers of existing predisposition to AMD, and indeed we (Zhang) and others have performed genome-wide association studies (GWAS) to demonstrate that development and progression of AMD is strongly tied to a number of loci, such as the complement pathway genes complement factor H (CFH), C3, CFB, and CFI (10-14). Another highly significant AMD-associated region has been identified and corroborated extensively (15-18), and lies on chromosome 10 at 10q26. Unlike the complement loci associated with AMD, the 10q26 region encompasses three different genes, each of which could be associated with AMD. The protein encoded by one of the genes, *HTRA1*, is found in drusen deposits, and is therefore an attractive causal agent, though conflicting results have implicated both it and the adjacent *ARMS2* as being affected by the AMD-associated risk haplotypes in the region.

The *HTRA1/ARMS2* locus.

The significant AMD susceptibility locus at 10q26 contains three candidate target genes, *PLEKHA1*, *ARMS2* and *HTRA1* (**Fig. 2**). Many sequence variants have been found in the region, contained within a number of risk and protective haplotypes. The primary risk haplotype has been refined to likely encompass ~21.5kb of sequence and two of the target genes: *ARMS2*, a gene of unknown function, and *HTRA1*, a secreted serine protease (17, 18). The current literature presents contradictory evidence for which of these genes is the target of regulatory variants in the region. Early results suggested variants in the promoter region of *HTRA1* led to increased expression and were associated with an increased incidence of AMD (17, 18). Subsequent studies were unable to corroborate these results, and *ARMS2* was again implicated as potentially causative (19). More detailed investigation identified several new polymorphisms associated with a decrease in *ARMS2* transcript levels and increased likelihood of developing AMD (20). Still further analysis has posited that a combination of the two effects (loss of *ARMS2* transcript and increased *HTRA1* expression) is necessary (21). The Zhang lab has recently resequenced the *HTRA1/ARMS2* region, including several dozen kilobases flanking the primary risk haplotype, in a 200 case, 200 control cohort, and identified 22 variants within the primary locus along with an additional 45 in the larger associated region (21). Thorough characterization has suggested that the primary risk

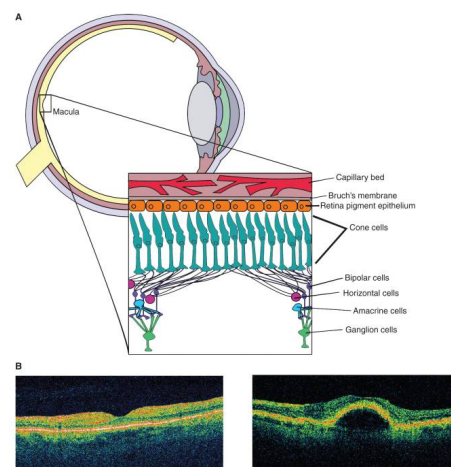


Figure 1. (A) Schematic of eye showing tissues affected by macular degeneration. (B) Optical coherence tomography images of a normal (left) and AMD retina showing RPE detachment (right).

and protective haplotypes can be distinguished by the identity of alleles present at four of these variants (**Fig. 2**). In order to identify the functional sites and targets of the risk haplotype, it is necessary to exhaustively assay all variants within the region, alone and in combination, and characterize their downstream effects.

Functional dissection of genomic variation with big DNA.

Nuclease-based genome engineering methods like CRISPR-Cas9 have come to the fore recently, but are less than ideal for high-throughput studies of multiple loci or those that require large genomic alterations, as they are capable of only one to a few edits at a time. Thus, testing many variants, especially in combination, is labor- and cost-prohibitive. Further, in diploid genomes, such one-change-at-a-time methods require extensive screening of clones in order to identify those with proper mutational phasing (scaling as 2^n clones for n mutations). Our method, in contrast, allows pairs or higher order combinations of variants of interest to be combined in cis in a single step, rather than randomly across to homologs. In general, such methods for large-scale ‘writing’ and wholesale replacement of large DNA sequences enable rapid high-throughput assessment of many genome variants either in isolation or combined in cis and importantly, many permutations can be assayed efficiently in parallel. The Boeke lab has been pioneering such technologies for the past several years, most notably in the design and synthesis of an entire eukaryotic genome with the Sc2.0 consortium (22). One such method for assembly of large DNA constructs is called “SWAP-In” and has been used to deliver segments of DNA ranging in length from 30,000 to 60,000 bp precisely to the desired location in the yeast genome by exploiting the endogenous homologous recombination (HR) mechanism in yeast, and moreover, this process can be iterated to produce synthetic yeast chromosomes of up to nearly 1 Mb (22-29). We are now applying these methods towards assembly of large mammalian loci and pathways called “Assemblons” because they are assembled by HR in the yeast cell. In particular, these assemblons are built by a specific instantiation of a homologous recombination-based method called eSWAP-In that we recently described (30). Importantly, these assemblons can be designed such that they allow combinatorial assembly of one or multiple variant segments interspersed with wild-type sequence segments.

In recent work (30), we have shown that it is possible to amplify a series of overlapping DNAs spanning the human *HPRT1* locus, and using “eSWAP-In” (extrachromosomal SWAP-In), rapidly assemble a 101 kb intact human *HPRT1* region in three sequential eSWAP-In steps. The resulting plasmid (essentially a designer BAC) was recovered in *E. coli* cells. The large insert was analyzed by restriction digests and shown to have the correct structure. The resulting insert DNA, along with flanking plasmid sequences was integrated into the X chromosome of mouse embryonic stem cells (ESCs), near the native murine *Hprt1* locus. DNA sequencing established that the entire human locus was delivered intact at the expected position. Immunoblotting established that the human gene was expressed in the ESCs.

Defining the impact of synthetic haplotypes.

Like most disease-associated variants identified by GWAS, most of the variants reported in the AMD-associated region in 10q26 lie in non-coding regions, either intergenic or intronic to the genes located there. Thus, the functions of these variants are likely regulatory in nature, either cis-acting at nearby genes or potentially trans-acting across the genome. We will therefore use gene expression measurements as the primary means for assessing functional impact of the AMD associated haplotypes. Gene expression effects will initially be local measurements of nearby genes (i.e. *HTRA1*, *ARMS2*, and *PLEKHA1*), since this seems to be the most likely explanation, but can be expanded to include transcriptome-wide measurements.

We will tackle this problem by designing and building an assemblon consisting of the entire *HTRA/ARMS2* locus and substantial swath of flanking DNA, encompassing roughly 120 kb. Once assembly is demonstrated, a library of variant assemblons can be rapidly produced using the same strategy. For example, suppose there are ten SNVs in linkage disequilibrium with the index risk SNV. It would be simple to prepare ten variants each bearing a single risk SNV, and 40 additional variants bearing all possible combinations of two SNVs. These could then be delivered to a suitable test cell, such as a retinal pigment epithelial (RPE) cell homozygous for the protective haplotype, replacing one of those “wild-type” alleles with a landing pad similar to that described by (30). This would allow the 50 engineered loci to replace one of the endogenous alleles in a single step. It would be important to “tag” the two alleles, e.g. with a fluorescent protein gene or a natural mRNA variant in order to distinguish the engineered alleles from the endogenous protective allele. Assuming the engineered allele is tagged, e.g. *HTRA1*-GFP and *ARMS2*-RFP, the readout of gene expression would be straightforward. Importantly, it is possible that this GWAS locus and perhaps many others is driven by not a single variant but by epistasis relationships among variants, and perhaps even expression effects on more than one gene. Indeed, in

the case of *HTRA1/ARMS2*, previous studies have implicated both genes separately as the target of AMD-associated regulatory variants (19, 20, 31, 32).

Results and Discussion

Assembly of a 119 kb ARMS2/HTRA locus.

With the aim of demonstrating the feasibility of our approach, we have completed a ‘first-draft’ assembly of the wild-type *HTRA1/ARMS2* locus. We chose a ~119kb region that encompasses both the *HTRA1* and *ARMS2* genes and the primary AMD risk haplotype identified previously (21) (**Fig. 2A**). The region was broken up into 34 segments averaging ~3.8 kilobases in length (2.1–4.1 kb), overlapping by an average of 370bp (**Fig. 2B–C**). Segments were obtained by amplification from a commercially available BAC clone containing the entire region. PCR amplicons were purified and co-transformed into yeast along with a linearized ‘assembly vector’ containing sequences overlapping the first and last amplicon (**Fig. 3B**). Importantly, assembly of the *HTRA1/ARMS2* assemblon was carried out in a single step rather than iterative assembly, meaning variant assembly will be easily completed in parallel. We identified complete assemblies by amplifying across the assembly junctions in an automated RT-PCR workflow (**Fig. 2D–E**), which were further fully sequence confirmed by whole-genome sequencing of the yeast strain (**Fig. 2F**).

The proposed combinatorial assembly of “synthetic haplotypes” will allow us to assess the regulatory impact of many variants within the risk region in isolation or combination, and thus in principle we will be able to narrow down and identify those variants likely to contribute to AMD development and progression. We can further characterize the functional impact of these sites by interrogating their effects on chromatin accessibility and transcription factor binding in relevant cell types.

The bigger picture.

GWAS studies have identified thousands of loci associated with thousands of diseases or other phenotypes (33). In over 70% of cases, the risk SNVs (single nucleotide variants) fall outside of coding exons, and are likely phenotypically associated through some unknown regulatory function. Generally, these risk SNVs are in linkage disequilibrium (LD) with a substantial number of other SNVs and sometimes other types of molecular variants such as indels, meaning they are typically inherited as part of a “haplotype block” of tens to hundreds of kilobases. This correlation between variants in the same region makes teasing apart causal variants difficult and unsatisfying as the set of risk-associated SNVs is rarely broken up by HR events occurring during meiosis. Given the number of potentially causative variants correlated with typical GWAS index variants, not to mention possible epistatic effects, current ‘top-down’ genome editing techniques leave much to be desired in attempts to identify the truly functional sites in these haplotypes.

We thus propose that GWAS hits can be more efficiently dissected by use of assemblon engineering and analysis of engineered human cell lines, as outlined above for AMD, as a general method, summarized in **Fig. 3**. An interesting alternative, in cases where the cell types of interest are difficult to access or difficult to transfect, is to instead use induced pluripotent stem cells (iPSCs) as the engineering platform, and then induce differentiation towards a relevant disease cell type. Extending this idea further would make use of embryonic stem cells (ESCs), either murine or human. Mouse ESCs have one very special advantage in that they can be developed completely into animals, from which any cell type can be analyzed or harvested. We and others have shown that human DNA sequences larger than 100 kb can be delivered to mESCs and not only support expression but development of viable animals. As shown for the alpha-globin locus by Higgs and his colleagues (34), human genes are even expressed only in the relevant cell types, albeit at a slightly lower level than the mouse gene. Related work by Duncan Odom, in which an entire human chromosome 21 was transferred to mESCs (35), showed that not only was the human chromosome segregated and replicated normally in the animal, but the genes were expressed in an accurately tissue specific manner. These two remarkable studies demonstrate that all or most of the regulatory sequences necessary for proper gene expression, including tissue- and organ-specificity, are highly conserved between mouse and human in a functional sense, implying that the transacting factors that recognize them can effectively reach across substantial species boundaries with great specificity.

Once risk and protective haplotypes are transferred to mouse ESCs, and either differentiated into an appropriate cell type, or converted to mice, the relevant transcript alterations should provide the information needed to develop a transcription-based assay for the risk haplotype. With this tool in hand, the variant assemblons can then be analyzed to fine-map the causative variant(s).

Methods

Assembly region definition

The region for assembly was chosen somewhat arbitrarily, while fulfilling a specific set of criteria. Namely, it had to encompass the previously identified primary AMD risk haplotype as well as the larger surrounding region resequenced by the Zhang lab (21). It also contained the entirety of the *HTRA1* and *ARMS2* genes, and a substantial portion of upstream and downstream sequence in order to capture as much potential regulatory sequence as possible. The ends of the region were further chosen to not interfere with any possible regulatory sites demarkated by histone modification or DNaseI hypersensitivity peaks as shown in the UCSC genome browser.

Primer design

Primers for segment amplification as well as assembly junction screening were chosen using an in-house developed algorithm. The algorithm was developed to select primers that are suitable for amplification of an arbitrary sequence from an arbitrary genome, so as to have minimal interference from duplicated or degenerate regions. Junction primers are designed to fulfill similar criteria, though with less strict parameters related to interference from other genome sequence.

Assembly segment preparation

Most assembly segments were amplified from a commercially available BAC (clone RP11-467N7 from the Chori BACPAC resources center, <https://bacpacresources.org>), using KAPA HiFi HotStart DNA polymerase (KAPA Biosystems). 32/34 amplicons were successfully amplified with default KAPA cycling conditions, and an annealing temperature of 65C and extension temperature of 68C. Amplicon 27 was successfully amplified with a 65C extension. Amplicon 34 required use of Q5 polymerase (New England Biosciences).

For assembly preparation, 3ul of the PCR reactions for segments 2-33 were pooled and purified using the Zymo Clean & Concentrator 5 kit. The entirety of the pooled products was transformed for assembly. Amplicons 1 and 34 were fused by PCR assembly to sequences corresponding to the linearized ends of the assembly vector to create segments overlapping the vector. These reactions were purified and added to the assembly transformation separately.

Assembly transformation

The pooled assembly segments and assembly vector were transformed using standard high efficiency yeast transformation methods. Briefly, cells are grown to mid-log phase before being collected and washed with lithium acetate. They are further with PEG, lithium acetate, carrier DNA, assembly DNA, and DMSO, before being collected again and resuspended in calcium chloride before plating. Cells were plated on SC media lacking leucine, as the assembly vector contains the LEU2 gene to enable selection.

Assembly clone screening

Initial screening for assembly clones was performed using high-throughput automated RT-PCR workflow developed in the Boeke lab. Briefly, colonies arising from the assembly transformation are selected and grown in liquid culture before isolating genomic DNA (also containing the assembly YAC DNA). Isolated genomic DNA is used as template in microscale RT-PCR reactions on a Roche 1536 lightcycler, and monitored for amplification. Shain and Clemens (36) introduced a method for robust evaluation of PCR amplification curves called maxRatio. The benefit of maxRatio is that two parameters are output to identify the exponential region of the amplification curve: functional cycle number and maxRatio = efficiency-1, which can be used to separate the positive and negative amplification reactions. An additional benefit is that positive reactions naturally cluster relatively far from the negative reactions, easing computational separation. An R package called qpcR (37) is used to implement the maxRatio method as well as melt curve analysis with iterative melting temperature identification and peak area calculation. R code has been written to parse the background-corrected fluorescence data, separate positive and negative reactions by maxRatio, assess consistency of maxRatio, and assign a binary positive/negative flag for each sample. Clones having positive reactions for all assembly junctions were then subjected to whole-genome sequencing of the yeast strain, mapping reads to a hybrid human and yeast reference genome.

Figure 2

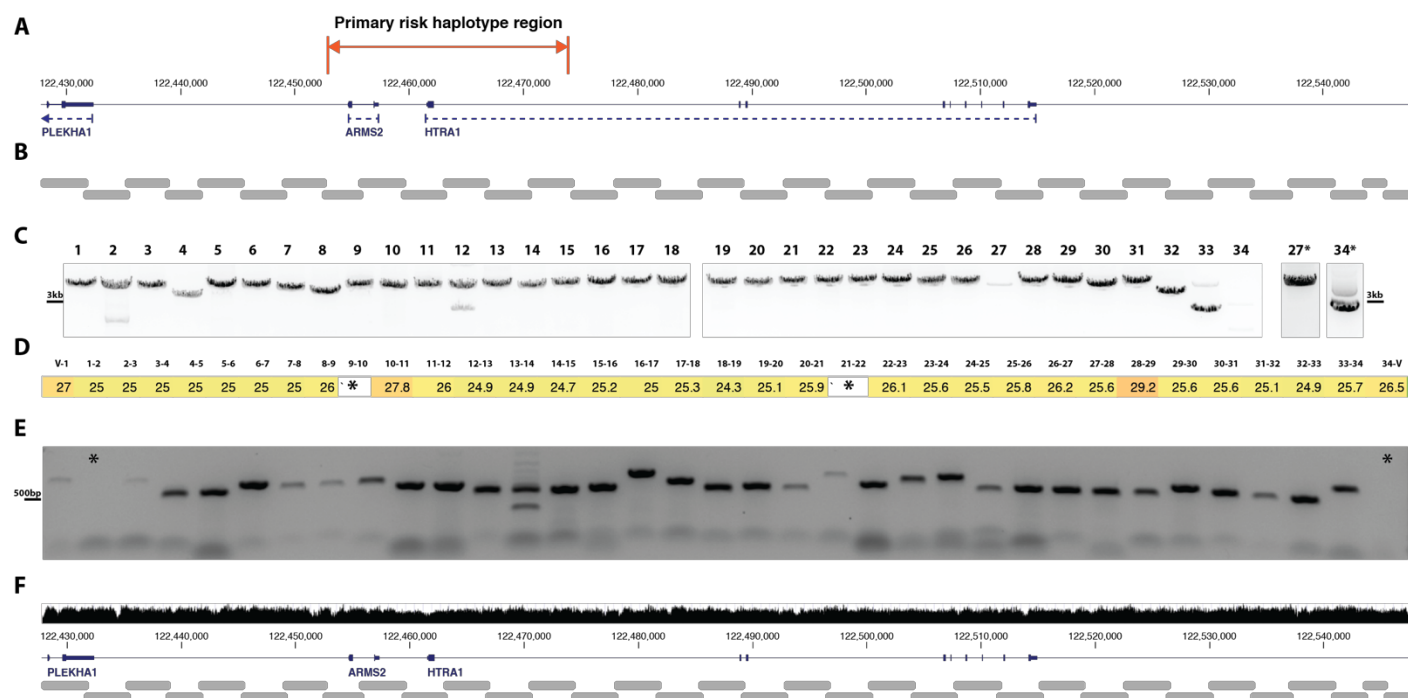


Figure 2. Assembly scheme for initial HTRA1/ARMS2 assemblon. (A) The genome locus surrounding HTRA1 and ARMS2 that is being assembled here. (B) Approximate size and position of the assembly segments designed by our automated platform. (C) PCR of assembly segments from a BAC clone covering the region. Segments 27 and 34 were amplified separately with alternative conditions, shown with *. (D) Example output from the automated RT-PCR junction PCR strategy showing crossing-point data and (E) manual junction PCR strategy for the same assemblon clone. * indicates junctions that did not amplify in the respective assay, but were ultimately present. (F) Sequence coverage track of the same assemblon clone in D and E obtained by WGS of the yeast strain.

Figure 3

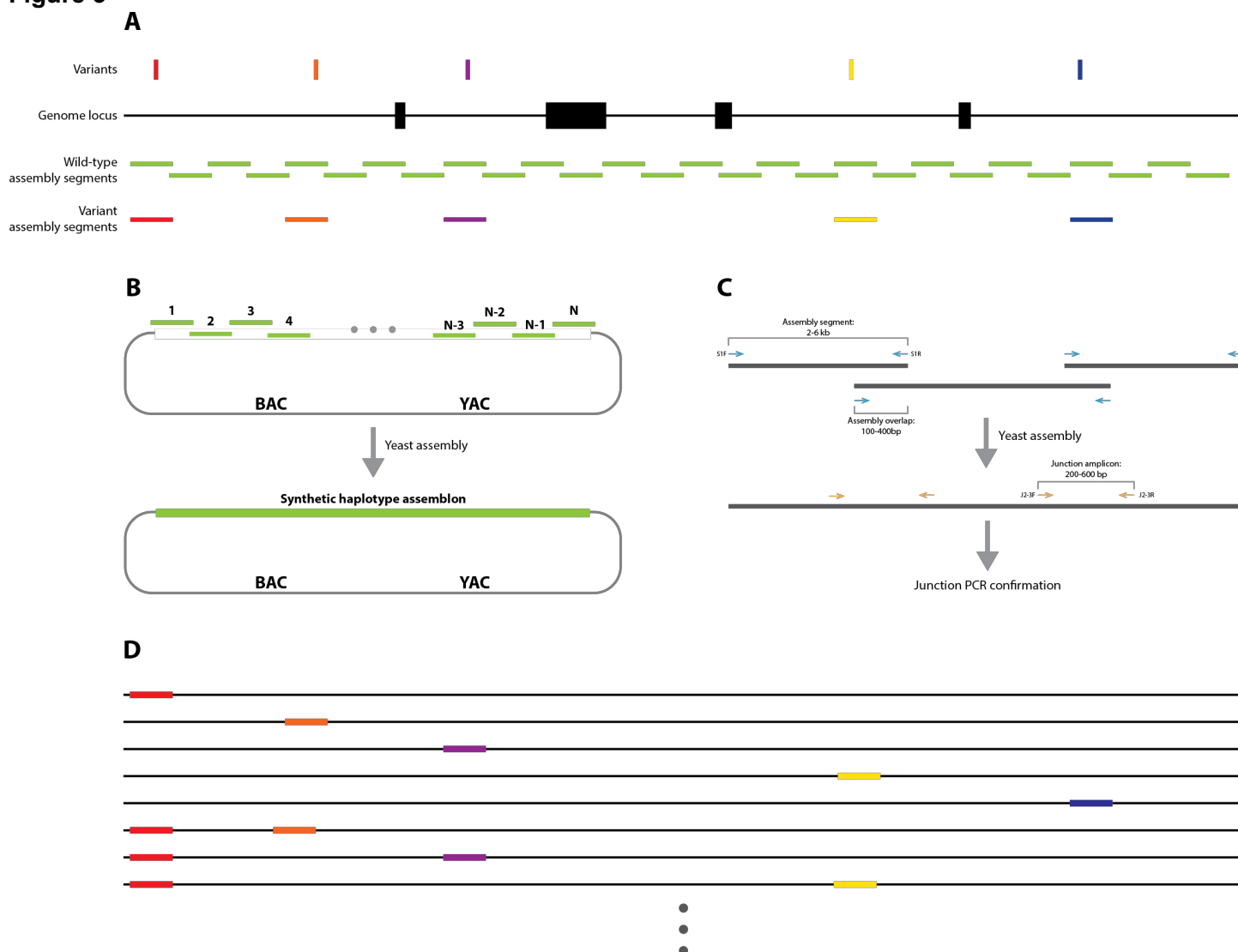


Figure 3. Generic scheme for interrogating disease-associated genetic variation with synthetic haplotypes. (A) A genome locus associated with some disease or phenotype, showing locations of associated variants, overlapping assembly segments, for wild-type and variant assembly. (B) Outline of the assembly method. Overlapping assembly segments are co-transformed into yeast along with a linear assembly vector, where they are efficiently assembled into a complete assemblon (C) Assembly segments are initially obtained as PCR amplicons or synthetic segments using 'segment primers'. Correct assembly structure is first screened by PCR across the assembly junctions with 'junction primers' (D) A subset of possible variant synthetic haplotypes that would be present in assembled variant library for the locus shown in A. These can be efficiently assembled and delivered in parallel, allowing high-throughput interrogation of which variant sites are functional.

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