Allelic Ratios and the Mutational Landscape Reveal Biologically Significant Heterozygous SNVs

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ABSTRACT The issue of heterozygosity continues to be a challenge in the analysis of genome sequences. In this article, we describe the use of allele ratios to distinguish biologically significant single-nucleotide variants from background noise. An application of this approach is the identification of lethal mutations in Caenorhabditis elegans essential genes, which must be maintained by the presence of a wild-type allele on a balancer. The h448 allele of let-504 is rescued by the duplication balancer sDp2. We readily identified the extent of the duplication when the percentage of read support for the lesion was between 70 and 80%. Examination of the EMS-induced changes throughout the genome revealed that these mutations exist in contiguous blocks. During early embryonic division in self-fertilizing C. elegans, alkylated guanines pair with thymines. As a result, EMS-induced changes become fixed as either G→A or C→T changes along the length of the chromosome. Thus, examination of the distribution of EMS-induced changes revealed the mutational and recombinational history of the chromosome, even generations later. We identified the mutational change responsible for the h448 mutation and sequenced PCR products for an additional four alleles, correlating let-504 with the DNA-coding region for an ortholog of a NFκB-activating protein, NKAP. Our results confirm that whole-genome sequencing is an efficient and inexpensive way of identifying nucleotide alterations responsible for lethal phenotypes and can be applied on a large scale to identify the molecular basis of essential genes.

FORWARD genetics in model organisms, which involves random mutation and isolation of a phenotype, laid the foundation for characterization of gene function. The bottleneck of this process lies in the identification of the molecular lesion responsible for the phenotype. The traditional approach for mutation identification involves three-factor mapping followed by several rounds of complementation testing using deficiencies and duplications. To reduce the number of candidate-coding regions, cosmids and fosmids are used to attempt to rescue the lethal phenotype (Janke et al. 1997; Simms and Baillie 2010). Finally, PCR analysis and DNA sequencing are used to confirm the molecular identity of the gene. This approach is laborious, time-consuming, and has very low throughput.

Technological advancements have provided methods to speed up the process of mutation identification. Recently, array comparative genomic hybridization (aCGH) was applied to identify single-nucleotide variations (SNVs) in the genomes of Saccharomyces cerevisiae (Gresham et al. 2006) and Caenorhabditis elegans (Maydan et al. 2009). This genome-wide approach allows rapid identification of a region of interest without mapping the mutation. Together with dense tiling arrays, aCGH could narrow down a SNV to within 10 bp (Maydan et al. 2009). However, this approach, which relies on sensitive hybridization, is unable to detect heterozygous mutations (Gresham et al. 2006; Maydan et al. 2009).

Whole-genome sequencing (WGS) is coming to the forefront as an attractive alternative for identifying molecular lesions (Cronn et al. 2008; Hobert 2010). Many researchers, including ourselves, have successfully identified SNVs and large genomic variations using WGS (Sarin et al. 2008, 2010; Shen et al. 2008; Doitsidou et al. 2010; Filbotte et al. 2010; Maydan et al. 2010; Rose et al. 2010). This approach has greatly facilitated the characterization of mutant phenotypes as well as many natural variants (Hillier et al. 2008). WGS is particularly

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useful for identifying hard-to-map alleles and genes that cannot be rescued by conventional transgenic fosmid or cosmid libraries. Nevertheless, almost all of the studies to date have focused on identifying homozygous mutations whereas identifying heterozygous mutations continues to be a challenge. Identifying heterozygous SNVs is an important step in genome analysis for understanding genomic variations and is generally relevant to many situations where allelic differences exist. In this report, we have developed a method for effectively identifying heterozygous mutations in C. elegans using lethal mutations as a model.

C. elegans is a self-fertilizing hermaphrodite whose genome becomes homozygous within a few generations. However, C. elegans with lethal mutations cannot be maintained as viable homozygous strains. This problem has been solved by the use of “balancers” to isolate, maintain, and characterize essential gene mutations (Edgley et al. 2006). In C. elegans, two commonly used classes of balancer are translocations (Rosenbluth and Baillie 1981) and duplications (Rose et al. 1984). Duplications that do not crossover with the normal chromosomes provide a third allele that is wild type and can rescue the lethal mutation, which is effectively maintained as a homozygote. In the case of the duplication-rescued strains, the allele frequency is 2:1 mutant: wild type. In this article, we describe the first use of Illumina sequencing to identify the DNA-coding region of an essential gene rescued by the duplication sd52 on chromosome I of C. elegans.

Materials and Methods

C. elegans strains

C. elegans is a self-fertilizing hermaphrodite and produces isogenic progeny within a few generations. Strains carrying homozygous mutations in let-504 rescued by the duplication sd52 were previously generated (Howell et al. 1987) and complementation tested (Howell and Rose 1990). The wild-type N2 (KR4848) is a derivative of a CGCN2 strain maintained in the Rose laboratory. The strain carrying a het-

erozygous tm4719 allele was kindly provided by S. Mitani (National Institute of Genetics, Mishima, Shizuoka, Japan). The strain KR5173, which carries the tm4719 allele balanced by hT2[bl-i-4(e937)] let-x(q782) qls48] I; III, was generated in this study. C. elegans strains were maintained at 20° as previously described (Brenner 1974).

Genomic DNA preparation

KR772 and KR4848 worms were grown on five 10-cm agar plates with Escherichia coli OP-50 until food was depleted (~5 days at 20°). Worms were collected and pelleted by washing the plate with M9 and centrifuged at 1500 × g for 1 min at 4°. The worm pellet was washed three times with M9, followed by 2–3 hr of incubation at room temperature to allow bacteria digestion. The worms were pelleted as before and finally resuspended in 0.5 ml of TE. The worms were frozen in −20° and lysed in lysis solution (50 μl 5% SDS, 2.5 μl 20 mg/ml Proteinase K) at 60° for 2 hr. Genomic DNA was purified using phenol/chloroform extraction and ethanol precipitation. The sample was treated with 4 μl of 5 mg/ml RNase A for 1 hr at 37°, followed by a second round of phenol/chloroform extraction and ethanol precipitation. Approximately 10 μg of DNA was sheared for 10 min using Sonic Dismembrator 550 (cup horn, Fisher Scientific) with a power setting of “7” for 30-sec pulses interspersed with 30 sec of cooling and analyzed on a 8% PAGE gel. A 180- to 220-bp DNA fraction was excised and eluted from the gel slice overnight at 4° in 300 μl of elution buffer [5:1, LoTE buffer (3 mM Tris–HCl, pH 7.5, 0.2 mM EDTA): 7.5 M ammonium acetate] and was purified using a Spin-X Filter Tube (Fisher Scientific) and by ethanol precipitation. The whole genome shotgun sequencing library was prepared using a modified paired-end protocol supplied by Illumina. This involved DNA end-repair, formation of 3’ A overhangs using Klenow fragment (3’–5’ exo minus), and ligation to Illumina PE adapters. Adapter-ligated products were purified on Qiaquick spin columns (Qiagen) and PCR-amplified using Phusion DNA polymerase for 10 cycles using the PE primer 1.0 and 2.0 (Illumina). PCR products of the desired size range were purified using a 8% PAGE gel. DNA quality and quantity was assessed using an Agilent DNA 1000 series II assay and Nanodrop 7500 spectrophotometer (Nanodrop), and DNA was subsequently diluted to 10 nM. The final concentration was confirmed using a Quant-iT dsDNA HS assay kit and Qubit fluorometer (Invitrogen). For sequencing, clusters were generated on the Illumina cluster station and paired end reads were generated using an Illumina GAII platform following the manufacturer’s instructions. Image analysis, base calling, and error calibration was performed using the V1.0 Illumina Genome Analyzer analysis pipeline.

Mutational density calculation

We collected coordinates of all the homozygous EMS changes. The genome was divided into overlapping bins of 2 Mbp, and we counted the number of EMS changes in each bin. The mutational rate for each bin was derived by dividing the number of EMS changes by the bin size. The value for each bin was collected and used to plot Figure 4.

Whole-genome sequencing and analysis

The genomic sequence of KR772 was aligned to the annotated sequence of C. elegans available at WormBase WS200 (http://www.wormbase.org) using BWA at the default setting (Li and Durbin 2009) and compared with the sequence of the wild-type strain KR4848. Genome analysis and visualization were done using Integrative Genomics Viewer (Robinson et al. 2011). The SNVs were called using VarScan (Koboldt et al. 2009) with the following parameters: --min-coverage 20–min-avg-qual 20–min-var-frequition 0.2–p-value 0.1. Candidate nucleotide differences for let-504 (h448) were further filtered to satisfy the following three criteria: (1) mutations that fall within genetic mapping range; (2) unique to mutant strain compared to the N2 strain (KR4848); and (3) allelic ratio falls between 60 and 90%.
**Fosmid transgenic rescue**

The fosmid WRM0614bH01 was injected into dpy-5 hermaphrodites to construct the transgenic strain BC8626, which carries myo-2::GFP physically linked to dpy-5(+). The strain BC8626 was mated to the fosmid, and individual GFP outcross hermaphrodites were isolated. The progeny of individual GFP hermaphrodites were examined for sterile GFP Unc animals, which could be (1) crossovers between the let-504 and dpy-5, (2) carrying both the fosmid and sDp2, or (3) fosmid rescues of let-504. In the third case, GFP Unc animals would continue to segregate Uncs (all with GFP) and arrest Dpy Uncs. The length of the GFP Uncs was measured, and gonadal indexing was used to test for a shift in the h448 phenotype.

**Complementation test**

We received a knockout allele of E01A2.4, tm4719, from S. Mitani (National Institute of Genetics). Animals homozygous for tm4719 are sterile as adults. From a mixed population, animals heterozygous for the tm4719 deletion allele were crossed with males heterozygous for hT2[bli-4(e937)] let-x (q782) qIs48 I; III, which carries an insertion of myo-2::GFP. GFP hermaphrodites were selected and tested for tm4719 by PCR. One hermaphrodite carrying tm4719 was used to establish a balanced strain. GFP tm4719 males from the above cross were individually crossed to a single KR772 hermaphrodite [sDp2; let-504(h448) dpy-5(e61) unc-13(e450)/let-504(h448) dpy-5(e61) unc-13(e450)]. The F1 progeny from these individual crosses were screened for sterile non-Dpy non-Unc non-GFP adults. If h448 fails to complement tm4719, we would expect sterile adults segregating in the outcross progeny. The tm4719 deletion in sterile non-Dpy non-Unc non-GFP adults was tested with the PCR primers.

**Results and Discussion**

**Mutant strain selection**

Identification of the molecular basis of lethal mutations is problematic for WGS because the animals cannot be grown as homozygotes in large amounts for DNA production. In selecting a mutant strain to characterize, we took into account how well-mapped the mutation was and the number of alleles that failed to complement it. We chose the h448 mutation that is in the essential gene let-504. The h448 allele is maintained as a homozygote by a rescuing wild-type allele on the duplication sDp2. The free (unattached to a normal chromosome) duplication, sDp2, covers ~7.3 Mbp of the left half of chromosome I (Howell et al. 1987). We chose a deleted interval, hDf7, that is in the sDp2-balanced region because it was mapped to a well-defined area of ~200 kbp and contained a small number of essential genes (Figure 1). Of the six complementation groups mapping within hDf7 (let-353, let-503, let-504, let-505, let-506, and let-507), let-504 had the most alleles (Table 1) (Johnsen et al. 2000). It was with these considerations in mind that the strain KR772, which carries let-504 (h448), was selected. Previous analysis showed that the phenotypes of the let-504 alleles ranged from larval arrest to sterile adults (Howell et al. 1987; Johnsen et al. 2000). Our strategy was to identify the let-504-coding region by inspection of the genome sequence in the hDf7 region and to validate its identity using DNA sequencing of PCR products from the additional alleles.

**Nonrandom distribution of G→A and C→T changes**

Genomic DNA of KR772 was prepared and sequenced using Illumina sequencing. For comparison, we prepared and sequenced the genome of KR4828, a Bristol wild-type (N2) strain from the Rose laboratory. In KR772, a total of 45,694,133 read pairs of 114 bp read length were generated. Approximately 87% of the reads were aligned to the annotated C. elegans genome (WS200) using BWA (Li and Durbin 2009). The number of reads provided ~80-fold coverage on average.

To better identify candidate mutations for let-504, we first analyzed the general mutational load of KR772. We compiled all the base-pair differences unique to KR772 using Varscan (Koboldt et al. 2009) (Supporting Information, Table S1). We observed 648 SNVs present with >90% read support and, of these, 55% (357) were either G→A or C→T changes (Figure S1), which are characteristic of EMS mutations (Bautz and Freese 1960; Greene et al. 2003). Even though the lethal mutation was induced using a relatively low dose of 15 mM EMS (Howell et al. 1987), compared to the 50-MM dose that is often used (Brenner 1974; Sulston and Hodgkin 1988; Filibotte et al. 2010; Sarin et al. 2010), there still appear to be a large number (357) of apparent EMS-induced changes across the whole genome.

We analyzed the positions of the homozygous SNVs in KR772 and found that these changes do not distribute evenly across the genome. Figure 2 shows the positions of G→A and C→T changes plotted separately along the chromosome. Surprisingly, EMS-induced mutations clustered in contiguous blocks of either G→A or C→T changes. In some cases, the blocks spanned the entire chromosome. We observed that on chromosomes II, V, and X, the changes are predominantly G→A, whereas those on chromosome I are predominantly C→T (Figure 2).

We explain our observations in the following way. The affected gametes of the EMS-treated parent will have some alkylated G’s. For simplicity, we consider only the alkylated G’s in the sperm (Figure 3A). In the first round of replication of embryonic cell division after fertilization, the alkylated G’s will mis-pair with T’s (Figure 3B). In the second round of replication, the T’s will pair with A’s (Figure 3C). This results in EMS mutations becoming fixed such that the alkylated G’s have been replaced with A’s. The mutational changes will be the same for the entire DNA strand. For example, the alkylated G’s from the plus strand in the gamete will be fixed as A’s, and the C’s (G’s from the minus strand) will be fixed as T’s (Figure 3C). Only one of these
possibilities will be segregated to the germ-line progenitor cell (P-cell lineage) and passed onto the next generation. Thus, the offspring will inherit either G→A changes or C→T changes along the entire chromosome for any one affected gamete. To test the generality of this observation, we examined available EMS-treated genomic sequences published by Flibotte et al. (2010). In the strains RB5002, VC1923, and VC1924, the G→A and C→T changes occur in long contiguous blocks similar to our observation (Figure S2).

We also observed both in our data and in data from Flibotte et al. (2010) that there is a shift from one block of EMS type to another within a chromosome (Figure 2 and Figure S2). For example, chromosome IV in KR772 has a stretch of G→A changes and shifts to a stretch of C→T changes. Similarly, we observed in chromosome V of VC1924 (as an example) where the EMS mutations shift from a block of C→T changes to a block of G→A changes and then back to C→T changes. The shift between blocks can be explained as a result of crossing over during meiosis and subsequent homozygosis in the self-progeny of the hermaphrodite. In these cases, the paternal chromosome may have contained only G→A changes and the maternal chromosome only C→T changes. Crossing over between the homologs during meiosis would result in a chromosome with a segment of G→A changes and another segment of C→T changes. In summary, WGS of EMS-treated strains provides a way of identifying the type of mutational change along large stretches of the chromosome.

In addition to differences in SNV types, we also observed differences in SNV density. A sparsely mutated chromosome would have a flatter slope whereas a densely mutated chromosome would have a steeper slope. We observed many chromosomes shift from a densely mutated segment to a sparsely mutated segment, or vice versa (Figure 2 and Figure S2). We calculated the density of EMS mutations (see Materials and Methods) and observed that a lower density of mutation averaged about two SNVs per mega base pair and a higher density of mutation averaged between four and six SNVs per mega base pair (Figure 4). The shift in mutational density along the chromosome is likely a result of meiotic crossing over between the paternal and maternal chromosome. If so, the mutational frequency in sperm differs from the frequency in oocytes. There is evidence for this difference in other species. In Drosophila, EMS is more effective when fed to males than to females (Lewis and Bacher 1968), suggesting that sperms have a higher mutational frequency. More effective repair mechanisms and an increased cytosolic volume in the oocyte that may act as a sink for alkylating agents could result in a lower mutation frequency. Thus, it is possible that hermaphrodite sperm are more sensitive than oocytes to mutation.

Identification of heterozygosity using allelic ratio

The left half of chromosome I has notably fewer homozygous mutations than the rest of the chromosome. We predicted that the SNVs in that region would have <90% read support due to heterozygosity. We counted the number of SNVs as a function of their allelic ratio (Figure 5). For a typical chromosome, most of the SNVs fall within a 90–100% allelic ratio (e.g., the green line in Figure 5). However, in chromosome I, we observed a bi-modal distribution of the SNVs, with one peak at 70–80% and another peak at 90–100% (black line in Figure 5). Nearly all of the SNVs in the 70–80% category are located in the sDp2 region (blue dashed line in Figure 5) whereas SNVs outside of the sDp2 region are within 90–100% (red dashed line in Figure 5). We conclude that the EMS-induced mutations are homozygous along chromosome I homologs and differ from the wild-type alleles on the duplication, resulting in an allelic ratio in the range of 70–80%.

Table 1 Alleles of let-504

| Strain | Allele | Mutagen | Genotype | Arrest stage |
|--------|--------|---------|----------|-------------|
| KR456  | h137   | EMS     | sDp2; let-504 (h137) dpy-5 (e61) unc-13 (e450) | Sterile adult |
| KR661  | h327   | Gamma radiation | sDp2; let-504 (h327) dpy-5 (e61) unc-13 (e450) | L2/L3 |
| KR772  | h448   | EMS     | sDp2; let-504 (h448) dpy-5 (e61) unc-13 (e450) | Sterile adult |
| KR1506 | h844   | EMS     | sDp2; let-504 (h844) dpy-5 (e61) unc-13 (e450) | L3 |
| KR1541 | h888   | EMS     | hT1; let-504 (h888) dpy-5 (e61) unc-13 (e450) | L2/L3 |
We reasoned that we could use the allelic ratio to determine the extent of the duplication. We plotted the distribution of allelic ratios in 1-Mbp intervals along the chromosomes (Figure S3). Chromosomes I-right, II, III, IV, V, and the X are predominantly homozygous for the SNVs whereas the left half of chromosome I is predominantly non-homozygous for the SNVs (Figure S3). On chromosome I, homozygous SNVs emerge between 7 and 8 Mbp, indicating that the \(sDp2\) boundary falls within this interval. A detailed examination of the region (Figure 6) shows that homozygous mutations emerge between 7.2 and 7.3 Mbp, corresponding to the location of the right breakpoint of \(sDp2\).

**Application of EMS mutation blocks and allelic ratios to identify candidate mutations in the hDf7 region**

The \(hDf7\) region contains 62 predicted protein-coding sequences. Of these predicted coding sequences, 43 are potentially essential genes by the observation of arrest phenotypes after RNA interference treatment (Fraser et al. 2000; Sonnichsen et al. 2005). Even with this shortened candidate list, identifying the molecular lesion for \(let-504\) among the candidates is a daunting task. WGS of a strain carrying one of the alleles therefore presents a viable approach to identifying the responsible mutation. To narrow down the candidates, we took advantage of the two genomic features discussed above: (1) EMS mutation types occur in contiguous blocks and (2) the allelic ratio is likely to be close to 70–80%. Our analysis of the EMS mutations showed that chromosome I is predominantly C→T changes, and thus we predicted that \(let-504\) (h448) is also likely to be a C→T change.

The strain KR772 carries flanking markers in addition to the lethal mutation \(let-504\) (h448), and we examined the sequence for these pre-existing mutations. One of the markers, \(dpy-5\) (e61), is situated in the \(sDp2\) region, and the duplication provides a wild-type copy of the \(e61\) mutant allele. At position 5,432,448 on chromosome I, 80% (45/56 reads) had an A whereas the remaining 20% of the reads had a C at this position, which is the nucleotide in the wild-type \(N2\) sequence. Our results are in agreement with the
The other marker, \textit{unc-13} (\textit{e450}), is situated outside the duplicated region, and thus we expected that 90–100% of the reads would correspond to the \textit{e450} allele. We observed that 96% (55/56 reads) had a T at position 7,435,169 in a gene encoding \textit{unc-13} (Ahmed et al. 1992), whereas in our N2 strain there is a C at this position. The change would result in a STOP codon replacing the normal glutamine (Q) residue in the 13th exon. Previously, \textit{e450} was known genetically to introduce a stop codon into \textit{unc-13} (Waterston 1981); however, this is the first report of the nucleotide change responsible for this allele.

Having demonstrated that we could correctly identify SNVs corresponding to known mutations, we set out to find the molecular lesion for \textit{let-504}. The \textit{let-504} gene was mapped previously to the \textit{hdF7} region (Johnsen et al. 2000). Genetic mapping data placed the left breakpoint of \textit{hdF7} to the right of \textit{unc-89}, and the right breakpoint to the left of \textit{anc-1}, thus positioning \textit{hdF7} between 4.10 and 4.32 Mbp on chromosome I (Figure 1). We used both manual and bioinformatics analysis looking for SNVs within the \textit{hdF7} region. Similar to the \textit{dpy-5} (\textit{e61}) mutation, we expected the SNV to have an allelic ratio close to 70–80%, and we discounted any SNV that was present in all the reads. For example, we found a single base-pair deletion in \textit{R12E2.1} that was present in all the reads, and we discounted it as a candidate mutation.

We identified three SNVs that satisfied our criteria (Table 2). A G→A change with 70% read support (35/50) was found in \textit{H31G24.3}. No other SNVs were found in \textit{H31G24.3}. Sanger sequencing revealed that the same mutation occurs upstream of \textit{H31G24.3} in all the \textit{let-504} alleles. Since the different alleles of \textit{let-504} have different arrest stages, this mutation is likely not the \textit{h448} mutation. A C→T mutation was found in the last intron of \textit{E01A2.1} and was supported by 75% of the reads (111/148). This mutation did not disrupt the coding
region nor any splice signals and is thus unlikely to be the cause of $h_{448}$. The third mutation was in $E01A2.4$ and caused a C→T change at position 4,132,191 in 67% of the reads (42/63). This SNV changed the third nucleotide of the first codon from ATG (methionine) to ATA (isoleucine) and effectively removed the start codon for $E01A2.4$. Given the allelic ratio (67%) and the fact that it is a C→T change, we propose that $E01A2.4$ is $let-504$.

Identification of $let-504$ as $E01A2.4$, a NFκB-activating protein ortholog

We PCR-amplified and sequenced the coding region of $E01A2.4$ from strains carrying the remaining four $let-504$ alleles (Table 1). Sanger sequencing revealed nonsense mutations in $E01A2.4$ for three of the alleles: $h_{137}$, $h_{844}$, and $h_{888}$ (Figure 7). $h_{888}$ changes the 41st codon from TGG (W) to TGA (STOP); $h_{844}$ changes the 151st codon from CAA (Q) to TAA (STOP); and $h_{137}$ changes the 358th codon from CAG (Q) to TAG (STOP). The fourth allele ($h_{327}$), which was generated with gamma radiation, did not contain a SNV in the coding region of $E01A2.4$ and may be a complex mutation not detectable by PCR amplification and sequencing.

The lethal phenotypes of $let-504$ alleles correlate with the positions of the alleles in $E01A2.4$. Using gonadal indexing, we determined that $h_{888}$ and $h_{844}$, which remove most of the protein sequence, have the most severe phenotypes, arresting at the mid-larval stage (Figure S4). The milder phenotype, that of sterile adults, was seen in $h_{137}$ and $h_{448}$ (Figure S4). The $h_{137}$ allele truncates the protein close to the end whereas the $h_{448}$ allele has a mutated start codon. The milder phenotype of $h_{137}$ may result from readthrough of the stop codon, and in the case of $h_{448}$ might be due to the use of an alternative start codon that allows a truncated protein to be made. However, we observed that the $tm_{4719}$ allele, which removes amino acids 257–404 in $E01A2.4$, also produces sterile adult animals. Thus, it is possible that the first 256 amino acids contain information required for progression beyond the mid-larval stage.

To further confirm that $let-504$ is $E01A2.4$, we carried out transgenic fosmid rescue and complementation test experiments. The transgenic rescue experiment was done by crossing $let-504$ ($h_{448}$) animals to a strain carrying the transgenic fosmid WRM0614bH01, which contains $E01A2.4$. However, we were not able to observe rescue. This could be for many

**Figure 5** Allelic ratio in KR772 for the whole chromosome I, whole chromosome III, part of chromosome I under sDp2, and part of chromosome I not under sDp2. Allelic ratio is presented as the percentage of reads that show SNV at a particular nucleotide position. In the sDp2 region, the peak at 70–80% represents mutations homozygous in the homologs with a wild-type allele in sDp2.

**Figure 6** Chromosomal I region between 6 and 9 Mbp. The blue bars represent nonhomozygous SNVs, and the red bars represent homozygous SNVs. An SNV with an allelic ratio between 40 and 89% is considered as nonhomozygous. An SNV with allelic ratio ≥90% is considered as homozygous. A nonhomozygous mutation first occurred at 7.3 Mbp.
reasons. Because of the sterility phenotype and morphological defects in the gonad, it is possible that E01A2.4 expression is required in the germ line, where expression of transgenic fosmid may be suppressed. In the absence of a rescuing construct, we carried out a complementation test with the tm4719 allele. We predicted that worms carrying both h448 and tm4719 alleles would arrest as sterile adults if let-504 is E01A2.4. Thus, we constructed h448/tm4719 animals (see Materials and Methods) and examined 83 sterile adults from h448 mothers, which also carried the tm4719 allele. Thus, h448 failed to complement tm4719. We conclude that E01A2.4 is the coding region for the essential gene let-504.

BLAST searches with the E01A2.4 protein sequence show sequence similarity to the human gene NFκB-activating protein (NKAP) with 30% identity at the protein level. In humans, NKAP is a transcriptional repressor that associates with the NOTCH corepressor complex and is required for T-cell development (Pajerowski et al. 2009).

Concluding remarks

The issue of heterozygosity continues to be a challenge in genome sequence analysis. Here, we have demonstrated that heterozygous SNVs can be identified effectively using information from the mutational landscape and allelic ratios. An application of our approach is the identification of lethal mutations in essential genes. About 3000–5000 of the 20,000 genes in C. elegans are estimated to be essential for development and survival. Over the past 25 years, thousands of lethal alleles corresponding to >500 essential genes have been isolated. These mutations are maintained as heterozygous mutations using translocations and duplications. In the case of the duplication sDp2, 237 essential genes have been mapped genetically. Correlating coding regions to these lethal mutations has been slow and laborious. We have shown that WGS is a time-efficient and cost-effective way for further characterizing essential genes. Our approach is also applicable to situations in which the heterozygous mutations exist in a 1:1 allelic ratio (m/+). With deep enough coverage, a 50% SNV allelic ratio will stand out against the statistical noise of the sequencing methodology. The coverage required will depend upon the read length and the sequencing methodology and can be calculated for specific situations. In addition, we have provided a better molecular understanding of EMS mutation fixation, which may be a useful tool for identifying alkylating agent-induced lesions in C. elegans and other model organisms. The fact that a particular type of EMS mutation occurs in contiguous blocks reduces the number of non-informative changes and provides prediction with regard to mutation type. The approach taken here is readily applicable to the

| Chromosome | Position | Reference | Mutation | Reference read support | Mutation read support | Allelic ratio (%) | Genomic environment |
|------------|----------|-----------|----------|------------------------|-----------------------|------------------|---------------------|
| I          | 4116834  | G         | A        | 15                     | 35                    | 70               | 18 bp upstream of H31G24.3 |
| I          | 4132191  | C         | T        | 20                     | 41                    | 67.21            | First exon of E01A2.4 |
| I          | 4142061  | C         | T        | 37                     | 111                   | 75               | Last intron of E01A2.1 |

Figure 7 Location of let-504 alleles. The changes underneath the allele name indicate amino acid changes.
rest of the lethal collection as well as to other phenotypes. As essential genes often encode highly conserved proteins that act either in cell maintenance or in a developmentally critical pathway, identification of the coding regions corresponding to the mutant collection will greatly increase both available genetic resources and information about gene function. Identification of the molecular basis of these genes is of value for both our understanding of animal biology and the study of human disease.

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Allelic Ratios and the Mutational Landscape Reveal Biologically Significant Heterozygous SNVs

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Figure S1  The distribution of single nucleotide variations (SNVs) in KR772.
Figure S2  Non-random distribution of EMS induced changes are seen in RB5002, VC1923, and VC1924. The X-axis represents the length of chromosomes. The Y-axis indicates each SNV ID. The data used to generate this figure was from the supplementary table of (FUBOTTE et al. 2010).
Number of SNVs per Mbp on ChrI

Number of SNVs per Mbp on ChrII

Number of SNVs per Mbp on ChrIII

Number of SNVs per Mbp on ChrIV

Number of SNVs per Mbp on ChrV

Number of SNVs per Mbp on ChrVI
Figure S3  Distribution of homozygous and non-homozygous SNVs per 1Mbp window in each chromosome. The blue bars represent non-homozygous SNVs and the red bars represent homozygous SNVs. A SNV with allelic ratio between 40% and 89% are considered as non-homozygous. A SNV with allelic ratio 90% or above are considered as homozygous.
**Figure S4**  DAPI staining and DIC images for h888, h844, h137, and h448. The yellow line outlines the gonad. The h888 and h844 alleles show the more severe phenotype where the gonadal arms have yet to turn. The h137 and h448 alleles show milder phenotype where the gonadal arms have fully turned.
Table S1  All SNVs present in KR772 with more than 20% read support.

Table S1 is available for download at http://www.genetics.org/content/suppl/2012/01/20/genetics.111.137208.DC1 as an excel file.