A rich TILLING resource for studying gene function in *Brassica rapa*

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Abstract

**Background:** The Brassicaceae family includes the model plant *Arabidopsis thaliana* as well as a number of agronomically important species such as oilseed crops (in particular *Brassica napus*, *B. juncea* and *B. rapa*) and vegetables (e.g. *B. rapa* and *B. oleracea*). Separated by only 10-20 million years, *Brassica* species and *Arabidopsis thaliana* are closely related, and it is expected that knowledge obtained relating to *Arabidopsis* growth and development can be translated into Brassicas for crop improvement. Moreover, certain aspects of plant development are sufficiently different between *Brassica* and *Arabidopsis* to warrant studies to be carried out directly in the crop species. However, mutating individual genes in the amphidiploid Brassicas such as *B. napus* and *B. juncea* may, on the other hand, not give rise to expected phenotypes as the genomes of these species can contain up to six orthologues per single-copy *Arabidopsis* gene. In order to elucidate and possibly exploit the function of redundant genes for oilseed rape crop improvement, it may therefore be more efficient to study the effects in one of the diploid *Brassica* species such as *B. rapa*. Moreover, the ongoing sequencing of the *B. rapa* genome makes this species a highly attractive model for *Brassica* research and genetic resource development.

**Results:** Seeds from the diploid *Brassica* A genome species, *B. rapa* were treated with ethyl methane sulfonate (EMS) to produce a TILLING (Targeting Induced Local Lesions In Genomes) population for reverse genetics studies. We used the *B. rapa* genotype, R-o-18, which has a similar developmental ontogeny to an oilseed rape crop. Hence this resource is expected to be well suited for studying traits with relevance to yield and quality of oilseed rape. DNA was isolated from a total of 9,216 M₂ plants and pooled to form the basis of the TILLING platform. Analysis of six genes revealed a high level of mutations with a density of about one per 60 kb. This analysis also demonstrated that screening a 1 kb amplicon in just one third of the population (3072 M₂ plants) will provide an average of 68 mutations and a 97% probability of obtaining a stop-codon mutation resulting in a truncated protein. We furthermore calculated that each plant contains on average ~10,000 mutations and due to the large number of plants, it is predicted that mutations in approximately half of the GC base pairs in the genome exist within this population.

**Conclusions:** We have developed the first EMS TILLING resource in the diploid *Brassica* species, *B. rapa*. The mutation density in this population is ~1 per 60 kb, which makes it the most densely mutated diploid organism for which a TILLING population has been published. This resource is publicly available through the RevGenUK reverse genetics platform http://revgenuk.jic.ac.uk.

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Background

The advent of high-throughput sequencing technologies, vast genomic databases and increasingly powerful genetic tools has had a huge impact on the development of our understanding of the biochemical and developmental networks regulating the multitude of genetic and physiological processes in plants [1]. Insight from studies in the model species, *Arabidopsis thaliana*, is increasingly facilitating our ability to elucidate and beneficially exploit key regulatory processes in relevant crop species. The last decade has seen the development of a number of large-scale ‘Reverse Genetics’ tools to study the effects of mutations in genes for which the sequence is known. These tools include T-DNA insertion [2], TILLING (Targeting Induced Local Lesions In Genomes) [3] and RNAi technologies [4-6].

TILLING is a reverse genetics tool, which was originally developed for *Arabidopsis* [7] and has subsequently been successfully employed in other plant species as well as animal species (eg. [8-17]). For plants, large mutant populations are generated by the treatment of seed or pollen with a chemical mutagen - most commonly ethyl methane sulfonate (EMS) - that can induce point mutations at a very high density, sufficient to establish a series of allelic mutations in all genes. Amplified sequences are then screened using established high throughput SNP discovery methods.

*Brassica napus* (oilseed rape) is an amphidiploid species containing two diploid genomes originating from a cross between the diploid *Brassica* species, *B. rapa* and *B. oleracea*. Whereas TILLING populations have been described for *B. napus* and *B. oleracea* [13,15], such a resource has not yet been reported for *B. rapa*.

EMS is a mutagenic, teratogenic and possibly carcinogenic organic compound and it is the mutagen of choice for the development of plant TILLING populations [7-15]. It produces random mutations in genetic material by nucleotide substitution; primarily by alkylation on the O6 position of guanine leading to GC→AT transition changes.

Here we describe the development of a TILLING population in *B. rapa* genotype R-o-18 [18]. The effect of EMS on plant growth and fertility in the M1 and M2 generations is described. Based on the screening of six genes located on different chromosomes, we calculated a mutation density of ~1 per 60 kb and a 97% probability of identifying a stop-codon mutation in a standard screen of 3072 M2 plants. This resource therefore comprises an attractive tool for researchers having interests in plant development and especially with regard to phenotypic traits related to improvement of oilseed rape and other crops.

Results and Discussion

Choice of model plant

R-o-18 (Figure 1) is an inbred line of the *Brassica rapa* subsp. *trilocularis* (Yellow Sarson) with transparent seed coat [18] closely related to *B. rapa* oilseed crops grown in Pakistan [19]. It was favoured as the genetic system for the TILLING population described here due to the following features: 1) it is diploid, which simplifies the genetics and reduces the potential for genetic redundancy, 2) *B. rapa* genome sequencing is ongoing and sequence information on the target gene is a prerequisite for reverse genetics, 3) R-o-18 is self fertile and produces a large number of seeds per plant (this is in contrast to several other *B. rapa* varieties such as the subspecies *chinensis* and *pekinensis*) and 4) the similarity in growth pattern to both the model plant *Arabidopsis thaliana* and oilseed rape varieties makes it an attractive system for fundamental research as well as studies related to yield and quality traits in *B. napus*. Such traits include seedling establishment, plant architecture, seed oil composition, fruit development, nutrient and water use efficiency and disease resistance.

Optimising mutagen dosage

Under optimal mutagenesis conditions, individuals of an EMS mutant population carry a high mutation load but remain vigorous and fertile. It is important, therefore, to determine the level of mutagen treatment necessary to achieve the maximal mutation load. We carried out incubations of R-o-18 seeds with EMS concentrations ranging from 0 to 1% (Figure 2).

EMS is a highly volatile and unstable compound and we have previously observed variation in the activity between batches. In order to avoid having to repeat the optimisation experiment with a different EMS batch after the titration, we used the same fresh batch of EMS to treat 5,000 seeds for each concentration in the 0.2-0.5% range (expected to be most relevant for producing the population based on previous experience) and 200 seeds at the concentrations outside this range.

After incubation and washes, the seeds were sown in soil and kept at 7°C for six days. Following a further six days in the glass house, germination frequency was established. Germination was hardly affected by treatments up to 0.3%. However, at 0.4% a marked decrease was observed and at 1% EMS none of the seeds germinated, indicating that the EMS treatment had been effective.

For *Medicago truncatula* mutagenesis, it was previously reported that an EMS concentration at the point where germination begins to become compromised is optimal for obtaining a large mutation load while maintaining vigorous and fertile plants [20]. For the *B. rapa* population we therefore decided to use seedlings derived...
from the 0.3% and 0.4% EMS treatments for population development, and the resulting two populations will subsequently be referred to as the ‘0.3% population’ and the ‘0.4% population’.

Although, only seedlings from the 0.3% and 0.4% EMS-treated seeds were used to make up the M1 generation of the mutant population, a subset of the germinated seedlings from the remaining concentrations in the titration experiment were allowed to grow on. As shown in Figure 2b and 2c, the higher concentrations of EMS also inhibited seedling establishment as well as plant growth.

Twenty M1 plants from each concentration were grown to maturity to establish the overall population fertility. The total number of viable seeds decreased gradually throughout the concentration range; the number of aborted seeds first increased as expected and then decreased at the higher concentrations probably due to effects on early ovule development (Figure 3a). The number of seeds per pod also decreased gradually with a severe reduction already apparent at 0.3% and 0.4% EMS (Figure 3b). The viability of the isolated seeds was established in a germination assay (Figure 3c). Despite a clear reduction in seed number at the low EMS concentration, no effect was observed on M2 seed viability until the 0.3% EMS level. This could be due to the lack of homozygosity at a sufficient number of loci. However, the huge variation in seed viability observed at the 0.3% EMS level, suggests that a high mutation frequency has been reached at this concentration. In total, we obtained M2 seeds from 3,464 and 1,564 M1 plants from the 0.3% and 0.4% populations, respectively.

M2 population and TILLING platform

When treating the seeds with the mutagen, a subset of cells in the shoot apical meristem of the embryo will carry the mutations on to the next generation. This provides an opportunity for multiple cell lineages to be
subject to a different spectrum of mutations, and it is therefore possible for gametes arising from different floral primordia to carry a distinct subset of mutant alleles [21].

Since *B. rapa* plants are larger, generate less seed and have a longer life cycle relative to, for example, *Arabidopsis thaliana* ecotypes Col-0 and L-er (5-6 months versus 6 weeks from seed to seed), it is desirable to minimise the number of plants necessary to build a useful resource. Under optimal conditions, we estimated that a sufficient number of the mutations would be recovered by using material from two *M2* plants from each of the ~5,000 *M1* plants assuming a similar number of progenitor cells as in *Arabidopsis*. Ten seeds from each *M1* individual were planted to increase the probability that at least two would germinate.

During growth of the *M2* population a number of phenotypes were observed; the percentage of *M2* families with albinos was 4.5% for both the 0.3% and 0.4% populations and we observed a plethora of morphological defects at developmental stages, ranging from seedlings to the fruit stage. Galleries of selected phenotypes are shown in Additional files 1, 2, 3 and 4.

**TILLING platform design**

For most *M2* families, 5-10 seeds germinated and in these cases we always took leaf tissue from the two most healthy-looking individuals and discarded the rest.
In this way, we obtained vigorous and mostly fertile plants, while expecting to maintain the high mutation level in a heterozygous state. The plants were subsequently bagged to prevent pollination between plants and M2 seeds were harvested.

Upon harvesting, we recorded the fertility and found that 9.6% of the M2 plants from the 0.3% population and 27.9% of the M2 plants from the 0.4% population failed to set seeds suggesting a higher mutation load in the 0.4% population.

DNA was isolated from the tissue, the concentration of DNA accurately determined and stocks normalised to ensure that DNA pools were balanced such that all individual lines were equally represented within the pools.

We used a standard one-dimensional pooling strategy where each M2 line is represented only once in a single pool, with each pool comprising DNA from eight M2 lines. Such a design is ideally suited to high throughput mutation detection. Specifically, eight-pools of 6,912 M2 plants originating from the 0.3% population were distributed in nine 96-well plates (DNA from 768 M2 plants per plate), whereas DNA pools from 2,304 M2 plants from the 0.4% population were distributed in three plates.

Mutations in genes of interest are detected by Cel1 digestion at a mismatched base pair [22,7]. For the B. rapa population described here, identification of digested fragments were carried out on an ABI3730 sequencer using fragment lengths of ~1 kb and a previously established protocol [14]. Individual M2 lines from pools identified as containing a mutant allele were subsequently sequenced in order to confirm the presence of the mutation, reveal its identity and to identify subsequent lines. Such a design is ideally suited to high throughput mutation detection. Specifically, eight-pools of 6,912 M2 plants originating from the 0.3% population were distributed in nine 96-well plates (DNA from 768 M2 plants per plate), whereas DNA pools from 2,304 M2 plants from the 0.4% population were distributed in three plates.

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EMS mutagenesis and the genetic code
When initiating a TILLING screen in a particular gene of interest, it is useful to analyse the coding region with reference to the genetic code. Only a subset of all combinations of amino acid changes are achievable when using EMS as a mutagen. Firstly, eight out of the 64 codons (12.5%) are unaffected by EMS-induced mutations because they do not contain guanine or cytosine. Secondly, out of the 96 positions that can be mutated (G→A or C→T) within the genetic code, 33 would not lead to an amino acid sequence change (silent mutations). Of the remaining 63 mutable positions of the genetic code, 58 would give rise to 26 amino acid substitutions (mis-sense mutations) and 5 would result in stop codons (nonsense mutations). Nine of the possible amino acid changes (corresponding to mutations at 21 out of 58 sites) result in chemically similar amino acids being incorporated which in many cases would be less likely to alter the function of the encoded protein significantly.

Mutations leading to premature stop codons are often desirable as they are expected to provide a dramatic reduction in gene function, especially when proximal to the 5’ end of the open reading frame. However, out of the 96 mutable positions, there are only five ways in which a stop codon can be obtained. These comprise the two glutamine codons (CAA and CAG), one of the six arginine codons (the C of CGA) and the tryptophan codon (TGG) for which G→A mutations at either position will generate a stop codon. The genetic code therefore has considerable robustness built-in, which minimises the potential biological effect of point mutations. It may therefore be beneficial to target the analysis of the gene of interest to a region in the sequence where it is possible to realise fully the potential of mutations that may reduce or abolish the activity of the encoded protein. To assist us in this analysis, we use the software package CODDLE [3], which is a programme designed to identify areas within the gene with highest probability of affecting gene function when mutated by EMS.

Amplicon selection
Selecting a suitable amplicon for mutation detection is a pre-requisite for establishing efficient and successful screens, and several considerations need to be taken into account: 1) In the large majority of cases, it will be advantageous to include as much coding sequence as possible and avoid intron or intragenic sequence. 2) Repetitive sequence may cause ‘Tag slippage’ which could delete or insert extra repeats. This will lead to artificial mismatches between wild type and ‘slippage’ strands, which may become substrates for the Cel1 nuclease. 3) As mentioned above, only a subset of codon-changes is likely to have dramatic effects on gene activity. It is therefore advisable to identify the region with most potential for generating stop codons and significant amino acid changes. 4) Finally, it is important to test for paralogue-specificity as up to three copies of each single-copy Arabidopsis gene may be present in the diploid Brassica genome [23,24]. When designing primers for the region of interest, it is therefore essential to verify that this primer set only amplifies the expected sequence before initiating the TILLING screen.

TILLING assays
We identified six genes that are located on different B. rapa chromosomes. These were expected to be orthologues of the Arabidopsis REPLUMLESS (RPL; At5 g02030) [25], INDEHISCENT (IND; At4 g00120) [26] and METHYLTRANSFERASE1 (MET1; At5 g49160) [27]
genes and were named BraA.RPL.a, BraA.RPL.b, BraA.RPL.c, BraA.IND.a, BraA.MET1.a and BraA.MET1.b, respectively, according to the accepted gene nomenclature system for the Brassica genus [28]. These B. rapa gene sequences were isolated using the Arabidopsis thaliana Integrated Database http://atidb.org as described in the Methods section.

We performed TILLING on ~1 kb amplicons from the six genes on the 0.3% population and for two genes on the 0.4% population. Based on the number of mutations identified for the individual genes, the TILLING populations were characterised with respect to mutation density and load (Table 1). The density was calculated at ~1 per 56 kb for the 0.3% population and ~1 per 67 kb for the 0.4% population after normalisation to the average 35% GC level for the B. rapa genome [29].

This density is the highest reported for a TILLING population in any plant or animal diploid species. Only in populations of tetraploid and hexaploid wheat [10,11] and the amphidiploid B. napus [13] have higher mutation densities been obtained.

Based on the stronger effect of the 0.4% EMS treatment on fertility, seed viability and general plant development compared to 0.3% EMS (Figure 2 and 3), it was unexpected that the number of mutations in the 0.4% population was not higher (Table 1). One explanation for this may be the increased number of homozygous loci observed in M2 plants of the 0.4% population (Table 2) and hence higher potential for lethality. We speculate that this difference in heterozygote: homozygote ratio may be due to the characteristic development in R-o-18 plants of two to four individual racemes derived from different progenitor cells. As a consequence, flowers on different racemes may have a different set of mutations, and since the whole plant is bagged, outcrossing between flowers of different progenitor origin is free to occur. With a decrease in fertility between the 0.3% and 0.4% populations, a higher number of 0.4% plants will have sterile racemes, thereby leading to higher incidents of self-pollination events. We believe this is reflected in the higher occurrence of homozygous plants in this population (Table 2).

### Table 1 Results from eight TILLING assays in the B. rapa mutant population.

| Gene name   | EMS (%) | Length (bp) | GC (%) | Mutations detected | Screened M₂ plants | Est. mutations per M₂ plant* | M₂ plants in population | Mutation density** | Expected mutations kb⁻¹*** |
|-------------|---------|-------------|--------|--------------------|--------------------|-------------------------------|--------------------------|----------------------|-----------------------------|
| BraA.RPL.a  | 0.3     | 1007        | 47     | 21                 | 768                | 13577 (10111)                 | 6912                     | 1/37 (1/49)         | 188 (164)                  |
| BraA.RPL.b  | 0.3     | 1072        | 45     | 149                | 4608               | 15082 (11730)                 | 6912                     | 1/33 (1/43)         | 209 (190)                  |
| BraA.RPL.c  | 0.3     | 1001        | 47     | 132                | 4608               | 14309 (10656)                 | 6912                     | 1/35 (1/47)         | 198 (173)                  |
| BraA.IND.a  | 0.3     | 1004        | 41     | 35                 | 3072               | 5674 (4844)                   | 6912                     | 1/88 (1/103)        | 78 (78)                    |
| BraA.MET1.a | 0.3     | 1104        | 48     | 94                 | 3072               | 13858 (10105)                 | 6912                     | 1/36 (1/49)         | 191 (164)                  |
| BraA.MET2.b | 0.3     | 1007        | 47     | 89                 | 3072               | 14385 (10712)                 | 6912                     | 1/35 (1/47)         | 199 (173)                  |

Relevant averages

12814 (9693)    1/44 (1/56)    177 (157)

| Gene name   | EMS (%) | Length (bp) | GC (%) | Mutations detected | Screened M₂ plants | Est. mutations per M₂ plant* | M₂ plants in population | Mutation density** | Expected mutations kb⁻¹*** |
|-------------|---------|-------------|--------|--------------------|--------------------|-------------------------------|--------------------------|----------------------|-----------------------------|
| BraA.RPL.b  | 0.4     | 1072        | 45     | 57                 | 2304               | 11539 (8975)                  | 2304                     | 1/43 (1/56)         | 53 (48)                    |
| BraA.RPL.c  | 0.4     | 1001        | 47     | 40                 | 2304               | 8672 (6458)                   | 2304                     | 1/358 (1/77)        | 40 (35)                    |

Relevant averages

10106 (7717)    1/51 (1/67)    47 (42)

* The estimated number of mutations per plant was calculated using an estimated genome size of 500 Mb for B. rapa [30]. Numbers in brackets were obtained after normalising to the average 35% GC content for the B. rapa genome [29].

** Mutation density based on estimated number of mutations per plant divided by the 500,000 kb genome. Numbers in brackets were obtained after normalising to the average 35% GC content for the B. rapa genome [29].

*** The number of expected mutations per kb in the population was obtained by extrapolating the number of detected mutations in a subset of the M₂ plants to the whole population.

Numbers in brackets were obtained after normalising to the average 41% GC content for B. rapa exons [29].
Using 500 Mbp as an approximate genome size for *B. rapa* [30], it was deduced by extrapolation that each plant contains close to 10,000 mutations (Table 1). One might expect this level of mutations to be lethal. However, as only 11% of the genome is coding sequence [29], we expect no more than 1,100 point mutations within exons of which about 700 will have the potential to cause amino acid changes and approximately 50 could introduce new stop codons. With the high heterozygotes: homozygote ratio (Table 2) and a high level of redundancy, it may therefore not be totally unexpected that it is possible to generate a large number of M2 plants with such a high mutation density.

The true mutation density is likely to be even higher than calculated here as only five of the total of 617 detected mutations were found within 100 bp from the fragment ends (Figure 4). The difficulty in detecting mutations close to the TILLING primers has been reported previously when LI-COR sequencers are used and 80 bp are now routinely omitted from both ends when calculating mutation densities using this technique [31]. Although a similar effect is observed with the ABI3730 sequencer used here, the mutation-density calculations are based on the actual numbers obtained in the assays and therefore represent an estimation of the minimal density.

Mutations in the *BraA.RPL* and the *BraA.IND.a* genes were asymmetrically distributed along the amplicons (Figure 4). This likely reflects the location of the 5’ primers in the *BraA.RPL* genes approximately 300 bp upstream of the start codon in an AT-rich non-coding region, necessary to ensure locus specificity. At the 3’ end, however, the primer was positioned within the first exon in an area of more mutable sites (higher GC content).

The high number of plants in this population further contributes to its potential value as an efficient reverse genetics resource. In a standard assay, 3,072 M2 plants are screened which corresponds to a third of the combined populations. From our experiments, an average of 68.2 mutations will be recovered when analysing a 1 kb fragment (41% GC on average in *B. rapa* exons [29]) with a 97% probability of obtaining a stop-codon mutation according to the formula:

\[
P = 1 - \left(1 - \frac{5}{96}\right)^{\text{number of mutations}}
\]

However, even if the desired mutations are not obtained in the first screen, there is still a further two thirds of the total number of lines available to be tested. In fact, extrapolation of the numbers obtained in the TILLING assays reported here shows that mutations in approximately half of the Gs or Cs in the *B. rapa* genome may exist in this population:

\[
\left(\frac{9,693 \text{ mutations per plant} \times 6,912 \text{ plants}}{1,000,000,000 \text{ bp} \times 0.35 \text{ GC bp}^{-1}}\right) \times 100\% = 48.4\%
\]

Bearing these numbers in mind, it is therefore surprising that no nonsense mutations were detected within the first 3,072 M2 plants for two out of the six amplicons tested here (*BraA.RPL.b* and *BraA.RPL.c* in Table 1). Screening an additional 1,536 M2 plants also did not result in any stop codon ‘hits’, whereas two nonsense mutations were detected for *BraA.RPL.b* after screening the 2,304 plants from the 0.4% population (Table 2).

It is unlikely that this discrepancy is due to lethality of nonsense mutations in these genes. Firstly, we have identified three closely related paralogues suggesting that these genes may function redundantly. Secondly,
mis-sense mutations in each individual do not have any detectable effect on plant development (data not shown).

These observations suggest that the lack of nonsense mutation detection in these amplicons is due to limitations of the detection method, which may be related to features in the DNA sequence.

**Gene Redundancy**

Although *B. rapa* is diploid, it is still a paleopolyploid having undergone an ancient triplication event [23,24,32,33]. Therefore one can expect to find up to three paralogous genes of each single-copy *Arabidopsis* gene in the *B. rapa* genome. Subfunctionalisation may have evolved in some cases. However, a high level of functional redundancy among the paralogues probably exists, and it may therefore be necessary to combine mutants to observe the desired effect. It is likely that this redundancy between paralogues allows *B. rapa* to harbour such a high mutation density compared to other diploid species. Another factor may be the high level of heterozygosity obtained especially in the 0.3% population, which to our knowledge is higher than for any other TILLING populations reported. The strategy of selecting healthy-looking *M₂* plants for the population will have contributed to this. One might predict that a significant number of *M₁* plants would be severely impaired in development due to a high level of homozygosity. In these cases it is recommended to carry out a backcross to wild type in the *M₂* generation to remove part of the background.

In accordance with the redundancy argument, we did not observe any developmental defects resulting from mutations in individual *BraA.*RPL paralogues. In contrast mis-sense and nonsense mutations in the *BraA.*IND.a gene, which is a single-copy gene in *B. rapa*, result in indehiscent phenotypes. With imminent availability of a complete *B. rapa* genome sequence expected in the very near future, this resource will have particular appeal since time-consuming gene isolation and design of parologue-specific primers will become a relatively straightforward informatic exercise.

### Linking mutation to phenotype

A classical backcrossing programme to remove the undesirable background mutation load is a prolonged procedure, which is expensive in both time and resources. This is especially true where a genus such as *Brassica* produces large plants with a relatively long generation time. Each backcross generation reduces the mutation load by 50%; so reducing the number of mutations from 10,000 to ten will take ten generations of backcrossing and genotyping (10,000 × 0.5₁₀ = 10).

As an alternative to embarking on such a programme, we instead assess the correlation between mutation and phenotype by comparing homozygous recessive mutants to heterozygotes and homozygous wild-type sibling plants in a segregating population for which the background mutations are the same. A 100% correlation between homozygous mutants and phenotype would lend strong support to the hypothesis that the mutation in the ‘TILLed’ gene is responsible for the phenotype. Moreover, we also aim to obtain allelic series of independent mutations in the same gene, where related phenotypic variants would strongly associate the phenotype with the gene, thereby avoiding the necessity for a lengthy backcrossing scheme. Detailed description on both of these approaches is provided in [34].

### Conclusions

Here we describe the development of a TILLING population in the *Brassica rapa* genotype R-o-18 suitable for reverse genetics studies. The high mutation density in this diploid species makes it an attractive genetic system for studying plant development and especially for obtaining mutations contributing to phenotypic traits related to crop improvement of oilseed rape. With the high mutation density, this resource will have particular appeal since time-consuming gene isolation and design of parologue-specific primers will become a relatively straightforward informatic exercise.

This population is publicly accessible and available via the RevGenUK reverse genetics platform http://revgen.nuk.jic.ac.uk.

### Methods

#### Mutagenesis and plant growth conditions

Seeds of the *Brassica rapa* genotype, R-o-18 (age 6-12 months since harvesting) were used in this work. For the negative control and 0.1%, 0.6%, 0.8% and 1% concentrations 200 seeds were treated in 10 ml solution in 50 ml Falcon tubes. For the 0.2%, 0.25%, 0.3%, 0.4% and 0.5% concentrations 5,000 seeds were treated in 250 ml (divided into 25 50 ml Falcon tubes for each of these concentrations). Seeds were soaked in 0.02% Tween 20 for 30 minutes prior to the addition of the EMS and incubated overnight (16 hours). 50 ml Falcon tubes with 200 seeds in each were turned end over end causing the seeds to tumble through the solution ensuring that all the seeds were exposed equally to the EMS without incurring too much physical damage. After treatment the seeds were washed ten times with 0.02% Tween 20 and then mixed with fine grade vermiculite to facilitate
their even distribution onto 348 mm × 220 mm seed trays. Seeds were sown at an approximate density of 200 seeds/tray and kept at 7°C with no light for six days before being transferred into the glasshouse at 18°C with 16 hours light. After a further six days the percentage of germination was recorded. These M1 seedlings were transplanted first into 24-well modules and then into 1L pots in John Innes no. 2 compost. The developing plants were bagged individually before they flowered using perforated bread bags (Packaging Company Unit, UK) to prevent cross-pollination, and allowed to proceed to maturity.

Dried pods were threshed and the M2 seeds from each line were deposited in the John Innes Centre seed-store (1.5°C, 7-10% relative humidity) to ensure their long-term viability. Ten seeds from each line were sown into 1L pots and transferred directly to the glasshouse. After germination the number of albino plants was recorded along with a range of other notable seedling and early leaf variant phenotypes. Each line was then thinned out to two ‘healthy’ plants per M2 line, leaf samples were taken for DNA isolation, and the plants were bagged individually and allowed to grow to maturity. Plants were harvested, threshed and M3 seed was placed in the seed-store.

DNA isolation, normalisation and pooling strategy
Leaf material was collected from each M2 plant selected for progression through to M3 seed and transferred directly into Qiagen racks on dry ice. DNA isolation was carried out using the DNeasy Plant 96 Qiagen Kit for 96 samples following the manufacturer’s instructions (Qiagen, UK). DNA concentrations were determined using PicoGreen (Molecular Probes, Invitrogen Corporation, Carlsbad, California, USA) against a universal DNA concentration standard on a Tecan Genios plate reader. All samples following the manufacturer’s instructions (Qiagen, UK). DNA concentrations were determined using PicoGreen (Molecular Probes, Invitrogen Corporation, Carlsbad, California, USA) against a universal DNA concentration standard on a Tecan Genios plate reader. All samples were normalised to 0.5 ng/μl (diluted in deionised water).

A simple one-dimensional eight-fold pooling strategy was employed using a Xiril liquid handling robot (Xiril AG, Hombrechtikon, Switzerland). Final DNA concentration in an eight-pool was 0.5 ng/μl, and 5 μl of eight-pool DNA were used in a TILLING reaction. This leads to a total of 2.5 ng DNA (~0.3 ng from each M2 line) in the TILLING reaction.

Primer design
Primers were designed using the Coddle (codons optimized to discover deleterious lesions; http://www.proweb.org/coddle) programme [3], combined with the PRIMER3 tool [35] to define the best amplicon for TILLING, aiming for a predicted primer T_m of 60-70°C. The primers used in this work are listed in Additional File 5. Coddle identifies areas within the gene which have the highest probability of affecting gene function when mutated by EMS, and scores possible mis-sense and nonsense changes.

TILLING
Mutant detection was carried out by Cel1 digestion followed by analysis on a capillary ABI3730 sequencer (Applied Biosystems, Foster City, California, USA) as described in [14].

Gene isolation and sequencing
The Arabidopsis thaliana Integrated Database (AtIDB - http://atidb.org) was used to blast the Arabidopsis genes, REPLIMLESS [25], INDEHISCENT [26] and METHYLTRANSFERASE1 [27] against a database of B. rapa BAC end sequences that have been mapped onto the Arabidopsis genome. BAC clones likely to contain the orthologous gene were identified based on synteny and primers were designed to sequence the genes directly on purified BAC DNA. No BACs were identified harbouring the Braa.MET1 genes. Instead these sequences were obtained from R-o-18 on a sequence homology approach based on a previous publication [36].

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Authors’ contributions
PS performed the EMS treatments and analysed the effect of titration. PS planned, organised and performed the planting, bagging, harvest and seed collection of the M₁ and M₂ generations and of tissue isolation for DNA extraction from M₂ plants. DB set up the TILLING platform and performed the screens. TG isolated and sequenced the BraA.IND gene and SA and GJK the BraA.MET7 gene. PS and LD analysed the data and wrote the manuscript. All authors read and approved the final manuscript.

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References
1. Østergaard L, Yanofsky MF. Establishing gene function by mutagenesis in Arabidopsis thaliana. Plant J 2004, 39(5):682-696.
2. Alonso JM, Stepanova AN, Leisse TJ, Kim CJ, Chen H, Shinn P, Stevenson DK, Zimmerman J, Barajas P, Cheuk R, et al. Genome-wide insertional mutagenesis of Arabidopsis thaliana. Science 2003, 301(5633):653-657.
3. Till BJ, Reynolds SH, Greene EA, Codomo CA, Enns LC, Johnson JE, Burtnet C, Odden AR, Young K, Taylor NE, et al. Large-scale discovery of induced point mutations with high-throughput TILLING. Genome Res 2003, 13(5):524-530.
4. Horiguchi G. RNA silencing in plants: a shortcut to functional analysis. Differentiation 2004, 72:65-73.
5. Schwab R, Oisowski S, Riester M, Warthmann N, Weigel D. Highly specific gene silencing by artificial microRNAs in Arabidopsis. Plant Cell 2006, 18:1121-1133.
6. Schwab R, Palanik JF, Riester M, Schormer C, Schmid M, Weigel D. Specific effects of MicroRNAs on the plant transcriptome. Developmental Cell 2005, 8:517-527.
7. McCullum CM, Comai L, Greene EA, Henikoff S. Targeting induced local lesions in genomes (TILLING) for plant functional genomics. Plant Physiology 2000, 123:439-442.
8. Perry JA, Wang TL, Welham TJ, Gardner S, Pike JM, Yoshida S, Parniske M. A TILLING reverse genetics tool and a web-accessible collection of mutants of the legume Lotus japonicus. Plant J 2003, 31:861-871.
9. Henikoff S, Till BJ, Comai L. TILLING. Traditional mutagenesis meets functional genomics. Plant Physiol 2004, 135(2):630-636.
10. Uauy C, Paraiso F, Colasuonno P, Tran RK, Talia H, Bernardi S, Comai L, Dubcovsky J. A modified TILLING approach to detect induced mutations in tetraploid and hexaploid wheat. BMC Plant Biology 2009, 9:115.
11. Slade AJ, Fuerstenberg SJ, Leeffler D, Steine MN, Faccott D. A reverse genetic, nontransgenic approach to wheat crop improvement by TILLING. Nat Biotechnol 2005, 23(1):75-81.
12. Till BJ, Reynolds SH, Weil C, Springer N, Burtner C, Young K, Bowers E, Codomo CA, Enns LC, Odden AR, et al. Discovery of induced point mutations in maize genes by TILLING. BMC Plant Biology 2004, 8:12.
13. Wang N, Wang Y, Tian F, King GI, Zhang C, Long Y, Shi L, Meng J. A functional genomics resource for Brassica napus: development of an EMS mutagenized population and discovery of FAE1 point mutations by TILLING. New Phytol 2008, 180(4):71-76.
14. Le Signor C, Savoie V, Aubert G, Verdier J, Nicolas M, Pagny G, Moussey F, Sanchez M, Baker D, Clarke J, et al. Optimizing TILLING populations for reverse genetics in Medicago truncatula. Plant Biotechnol J 2009, 7(5):430-441.
15. Himelblau E, Gilchrist EJ, Buono K, Bizzell C, Menter L, Vogelzang R, Osborn T, Amasino RM, Parkin IA, Haughn GW. Forward and reverse genetics of rapid-cycling Brassica oleracea. Theor Appl Genet 2009, 118(3):953-961.
16. Winkler S, Schwabedissen A, Backasch D, Bokel C, Siedel C, Bonisch S, Furthauer M, Kuhns A, Cobroes L, Brand M, et al. Target-selected mutant screen by TILLING in Drosophila. Genome Res 2005, 15(7):718-723.
17. Gilchrist EJ, O’Neil NJ, Rose AM, Zetka MC, Haughn GW. TILLING is an effective reverse genetics technique for Caenorhabditis elegans. BMC Genomics 2006, 7:262.
18. Ruhollha R, Higgins EE, Walsh JA, Lydiate DJ. Genetic control of broad-spectrum resistance to turnip mosaic virus in Brassica rapa (Chinese cabage). J General Virology 2007, 88:3177-3186.
19. Rana D, Boogaard van den T, O’Nell C, Hynes L, Bent E, Macpherson L, Park JY, Lim YM, Bancroft I. Conservation of the microstructure of genome segments in Brassica napus and its diploid relatives. Plant J 2004, 40:725-733.
20. Fournet RV, Cook DR. Production and characterization of diverse developmental mutants of Medicago truncatula. Plant Physiology 2000, 123:1387-1397.
21. Koornneef M. Classical mutagenesis in higher plants. Molecular Plant Biology. Oxford University Press:Gillmartin PM, Bowler C 2002, 1-11.
22. Olejárková CA, Bronson Mullins CR, Godwin AK, Yeung A. Mutation detection using a novel plant endonuclease. Nucleic Acids Res 1998, 26:4597-4602.
23. Lyak MA, Koch MA, Pecinka A, Schubert I. Chromosome triplication found across the tribe Brassicaceae. Genome Research 2005, 15:516-525.
24. Parkin IAP, Gulden SM, Sharpe AG, Lukens L, Trick M, Osborn TC, Lydiate DJ. Segmental structure of the Brassica napus genome based on comparative analysis with Arabidopsis thaliana. Genetics 2005, 171:765-781.
25. Roeder AH, Ferrándiz C, Yanofsky MF. The role of the REPLUMLESS homeodomain protein in patterning the Arabidopsis fruit. Current Biology 2003, 13:1630-1635.
26. Liangren SJ, Roeder AH, Kempin SA, Gremkzi K, Østergaard L, Guimil S, Reynolds DK, Yanofsky MF. Control of fruit patterning in Arabidopsis by INDEHISCENT. Cell 2004, 116:843-853.
27. Kato M, Miura A, Bender J, Jacobsen SE, Kukutanis T. Role of C and non-CG methylation in immobilization of transposons in Arabidopsis. Current Biology 2003, 13(5):421-426.
28. Østergaard L, King GJ. Standardized gene nomenclature for the Brassica genus. Plant Methods 2008, 4:10.
29. Trick M, Kwon S-J, Choi SR, Fraser F, Seumourour E, Drou N, Wang Z, Lee SY, Yang T-J, Mun J-H, et al. Complexity of genome evolution by segmental rearrangement in Brassica rapa revealed by sequence-level analysis. BMC Genomics 2009, 10:539.
30. Arumuganathan K, Earle ED. Nuclear DNA Content of Some Important Plant Species. Plant Mol Biol Rep 1991, 9:209-218.
31. Greene EA, Codomo CA, Taylor NE, Henikoff SJ, Till BJ, Reynolds SH, Enns LC, Burtnet C, Johnson JE, Odden AR, et al. Spectrum of chemically induced mutations from a large-scale reverse-genetic screen in Arabidopsis. Genetics 2000, 144(2):731-740.
32. Blanc G, Wolfe KH. Functional divergence of duplicated genes formed by polyploidy during Arabidopsis evolution. Plant Cell 2004, 16:1679-1691.
33. Blanc G, Wolfe KH. Widespread paleopolyploidy in model plant species inferred from age distributions of duplicate genes. Plant Cell 2004, 16:1667-1678.
34. Perry J, Welham T, Cheminant S, Parniske M, Wang T. TILLING. Lotus japonicus Handbook SpringerMárquez AJ, Stougaard J, Udvardi M, Parniske M, Spanik H, Saaalbach G, Webb J, Churazzz M, Márquez AJ 2005, 197-210.
35. Reznin S, Skaltsky H. Primer 3 on the WWW for general users and for biologist programmers. Bioinformatics Methods and Protocols: Methods in Molecular Biology Cambridge: Humana Press:Krawetz S, Misener S 2000, 365-386.
36. Fujimoto R, Sasaki T, Nishio T. Characterization of DNA methyltransferase genes in Brassica rapa. Genes Genet Syst 2006, 81:235-242.

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