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Introduction

Camelina sativa (hereafter, Camelina) is an oil seed crop of the Brassicaceae family that has attracted considerable attention because of its short growing season, and its productivity in geographic regions with limited rainfall and soil fertility (Iskandarov et al., 2014; Pilgeram et al., 2007; Zubr, 1997). Despite the increasing commercial interest in Camelina, a limitation to the wider use of its seed oil in biofuels, lubricants and food applications is its high content of polyunsaturated fatty acids, particularly linolenic acid (18:3), which accounts for 30%–40% of seed oil from most Camelina cultivars (Iskandarov et al., 2014). The high polyunsaturated content makes Camelina oil more susceptible to oxidation and food products derived from this oil more prone to rancidity (Frolich and Rice, 2005). To address this deficiency in Camelina oil quality, efforts have been directed at increasing the content of the more oxidatively stable oleic acid by suppression of FAD2 genes for the Δ12 oleic acid desaturase that converts oleic acid to linoleic acid (18:2) and linolenic acid (18:3) (Hutchreon et al., 2010; Kang et al., 2011; Nguyen et al., 2013). The result of this genetic modification is an increase in oleic acid content and corresponding decreases in polyunsaturated fatty acid (18:2 and 18:3) content of seed oils (Nguyen et al., 2013). Methods for FAD2 suppression in Camelina and other crops have included RNA interference (RNAi; Clemente and Cahoon, 2009; Graef et al., 2009; Jung et al., 2011; Nguyen et al., 2013), microRNAs (Belide et al., 2012), TALENs (Haun et al., 2014) and standard mutagenesis and selection (e.g. Kang et al., 2011; Pham et al., 2012; Thambugala et al., 2013; Wells et al., 2014).

The recent advent of the highly efficient and facile CRISPR/Cas9 system for gene editing (Cong et al., 2013; Jinek et al., 2012; Mali et al., 2013) in animals (Petersen and Niemann, 2015; Proudfoot et al., 2015) and plants (Weeks et al., 2015) offers the opportunity to determine whether the oil composition of Camelina seeds could be favourably altered by knocking out the activities of a few or all of the six fatty acid desaturase 2 (FAD2) genes present in the genome of this allohexaploid plant (Hutchreen et al., 2010; Kang et al., 2011). If successful, this strategy would increase oleic acid content and lower the content of linoleic acid, linolenic acid and other long-chain polyunsaturated fatty acids. Because the genomes of Arabidopsis and Camelina—and the FAD2 genes, in particular (Hutchreen et al., 2010; Kang et al., 2011; Nguyen et al., 2013)—share strong homology with each other, we designed sgRNA constructs for use in Camelina but tested them first in Arabidopsis to determine their efficacy in creating Cas9/sgRNA-mediated gene mutations and changing the fatty acid composition of seeds. This strategy assumed that Arabidopsis plants homozygous for FAD2 gene mutations could be obtained within two or three generations, whereas three or more generations would be required to inactivate most or all of the FAD2 genes in allohexaploid Camelina. The nuclear genome of Camelina, as well as that of other allohexaploids such as bread wheat, contains three separate subgenomes that, because they avoid intergenomic...
(homoeologous) recombination, behave as three distinct and separate diploid genomes (Comai, 2005; Feldman and Levy, 2012; Madlung and Wendel, 2013). While homoeologous recombination can be manipulated genetically in certain species, there is effectively no opportunity to replace a functional FAD2 gene with a defective allele from another subgenome through classical breeding techniques. In other words, homozygous or biallelic knockouts of FAD2 genes must be achieved independently for all three subgenomes if complete depletion of FAD2 enzyme activity is to be achieved.

In this report, we present evidence for efficient, multigenerational, knockout of FAD2 genes by the Cas9/sgRNA gene editing system in somatic and germ-line cells of Arabidopsis and Camelina leaves and seeds. We demonstrate in these initial experiments that such alterations lead to increases in oleic acid composition from ~16% to >50% and total monounsaturated fatty acid (18:1, 20:1, 22:1) from ~32% to >70% with concurrent decreases in linoleic and linolenic fatty acid content.

Results and discussion
FAD2 as a target for Cas9/sgRNA modification

The results presented here are the first report from a long-term project to significantly change the oil composition of Camelina seeds using the Cas9/sgRNA gene editing system to knock out the activity of FAD2, a key gene involved in the synthesis of polyunsaturated fatty acids. All three pairs of FAD2 genes in the allohexaploid, C. sativa, have all been shown to be active in developing seeds (Hutcheon et al., 2010; Kang et al., 2011; Nguyen et al., 2013). Thus, knocking out one or more of these genes should decrease the conversion of oleic acid to linoleic acid and linolenic acid (Figure S1). Because Camelina FAD2 genes share extremely high homology with their counterparts in the much easier and quicker to manipulate diploid plant, Arabidopsis thaliana (hereafter Arabidopsis) (Hutcheon et al., 2010; Kang et al., 2011; Nguyen et al., 2013), our strategy (discussed in detail below) has been to design Cas9 and sgRNA genes that often, but not always, target the same 20-nucleotide (nt) DNA sequence [23 nt, if the NGG protospacer adjacent motif (PAM) region is included] present in both Arabidopsis and Camelina genes.

As detailed below, seed fatty acid profiles in T2, T3 and T4 generations of transgenic Camelina were followed along with DNA sequence modifications in both leaf tissues and seeds for each generation.

For both Arabidopsis and Camelina, three 20-nt target sites for Cas9/sgRNA-mediated DNA cleavage of the FAD2 gene were selected. These 20-nt target sequences were oriented within each gene either in the same 5’ to 3’ direction as the reading frame of the gene (the ‘forward’ direction) or in the 5’ to 3’ direction on the opposite (reverse) strand of DNA. The location of each of these sites in the FAD2 genes of Camelina and Arabidopsis is shown in Figure 1a. Each of these sites was deliberately chosen to be located in the 5’ portion of each gene to ensure that gene disruptions altering the gene’s reading frame would produce a translation product lacking enzymatic activity. The sites were also chosen to contain a restriction enzyme cut site at the site of Cas9/ sgRNA-directed DNA cleavage three base pairs upstream of the PAM site. As described below, if Cas9/sgRNA activity results in DNA cleavage at this site and if the double-stranded DNA break (DSB) is repaired by the error-prone nonhomologous end-joining (NHEJ) DNA repair mechanism, the restriction enzyme site often will be destroyed. Thus, if the region containing this site is amplified using the appropriate PCR primers, amplicons containing nonmodified restriction enzyme sites will be cleaved, whereas

![Figure 1](image-url)
amplicons containing a destroyed restriction site will remain intact. Size analysis of the restriction enzyme-digested amplicons thus allows a simple qualitative and rapid means for detecting leaf or seed samples that contain (or do not contain) genes whose DNA sequence has been altered by the Cas9.sgRNA gene editing complex.

Drawings depicting the structures of the Cas9 gene and sgRNA gene constructs designed to recognize and cleave each of the three FAD2 gene target sites (Figure 1a) are provided in Figure 1b. The complete DNA sequence for each construct is provided in Supporting Information and Experimental procedures. In each construct, the Cas9 gene is driven by the cauliflower mosaic virus 35S gene promoter and terminated with the Agrobacterium tumefaciens Tnos gene termination sequence as described previously (Jiang et al., 2013, 2014). The sgRNA genes are driven by the Arabidopsis U6 promoter and followed by the termination region from the same gene. The construct targeting the Arabidopsis FAD2R1 site contained a hygromycin resistance gene to allow for rapid selection of transgenic seeds (Figure 1b). The three other constructs each contained a DsRed2 gene that allowed the use of a hand-held green fluorescent flashlight to easily detect and select red fluorescing transgenic T1 seeds (Lu and Kang, 2008; Nguyen et al., 2013) obtained from T0 plants produced by transformation of Camelina or Arabidopsis using the floral dip transformation technique. Importantly, as part of our strategy, only red fluorescing seeds were selected for further evaluation at each generation. Thus, assuming normal Mendelian inheritance of the T-DNA region containing the DsRed gene and the accompanying Cas9 and sgRNA genes, each plant analysed in this study had the potential to actively express the Cas9.sgRNA complex in most or all of its tissues. Important implications of this strategy are discussed below.

Extensive Cas9.sgRNA-mediated gene editing in leaf and seed tissues

Over 200 T1 Arabidopsis plants and 300 T1 Camelina plants that carried various Cas9 and sgRNA genes targeting FAD2 genes were generated and analysed during this study (Data Set S1). From these T1 plants, we produced large numbers of T2 and T3 plant progeny and seeds (Data Set S1) and limited samples of T4 seeds. Each plant generated in this study was given a unique name (e.g. AtFAD2R1 T1-10-8-7 is the 7th T3 progeny plant from the 8th T2 progeny plant from the 10th T1 Arabidopsis parent plant targeted for gene disruption at the R1 site of the FAD2 gene).

To speed the selection of plants carrying mutations in a targeted gene and to discard plants lacking Cas9.sgRNA-directed gene mutations, DNA was extracted from leaf tissue of transgenic T1 plants and used for PCR amplification of sgRNA target sites of interest using sets of primers listed in Table S1. As described above, each selected target site contained a restriction enzyme recognition sequence overlapping the expected Cas9.sgRNA cleavage site three base pairs upstream of the NGG PAM site, for easy detection of mutation events. As a result, if the Cas9/sgRNA complex created a DSB at the expected site and if an error occurred during DNA repair by the NHEJ mechanism, the restriction enzyme recognition site would be destroyed. In such cases, subjecting isolated DNA to digestion with the restriction enzyme appropriate to match the target site prior to PCR amplification of the target region would produce a nondigested, full-length, PCR product. Conversely, if no Cas9/sgRNA-directed mutation occurred at the site, the isolated DNA would be cleaved by the restriction enzyme and no PCR product would be visible on a standard ethidium bromide-stained agarose gel. An example of such a PCR/restriction enzyme (PCR/RE) analysis is shown in Figure S2 in which DNA from four different transgenic Arabidopsis plants displayed full-length, 322-bp PCR products, while DNA from a nontransgenic control plant produced the expected 244- and 78-bp restriction enzyme digestion fragments. Several DNA samples testing positive in the PCR/RE analyses were subjected to Sanger DNA sequencing of PCR-amplified target sites to determine the exact nature of the mutation. In most experiments, a few DNA samples tested negative in the PCR/RE analyses were also sequenced as controls and to confirm that they, indeed, were true negatives.

The expected ability of the Cas9/sgRNA system to cause gene mutations in the FAD2 genes of Arabidopsis and Camelina plants was confirmed by DNA sequencing. Sequence analyses of DNA from leaf and seed samples (Data Set S2) confirmed multiple mutations over multiple generations at each of the three target sites in each of the three different FAD2 gene types present, respectively, in the A, B and C subgenomes of the allohexaploid genome. In the overview presented in Data Set S2, DNA sequences obtained by Sanger sequencing are provided for 328 FAD2 gene target sites, 258 of which contained the kinds of short-length nucleotide insertions and deletions typical of NHEJ DNA repair that follows creation of DSBs by the Cas9/sgRNA complex. Interestingly, over half of these mutations were the result of a single-nucleotide insertion that was nearly always located at the predicted Cas9/sgRNA cut site 3 bp upstream of the PAM site. Among these single-nucleotide insertions, there was a marked preference for T (50%) and A (33%) nucleotide inserts over G (12%) and C (6%) nucleotide inserts. The summary at the bottom of Data Set S2 shows that approximately 19% of mutations involved nucleotide deletions over 3 bp in size and approximately 5% involved nucleotide insertions of 3 bp or more. Because all single-nucleotide insertions and most other insertions and deletions change the reading frame of the gene and because all target sites were deliberately chosen to be located in the 5’ region of the gene (Figure 1a), the vast majority of mutations created in the FAD2 genes are predicted to lead to gene knockout. Because of single-nucleotide polymorphisms (SNPs) between the three types of FAD2 genes found in the A, B and C subgenomes of Camelina (Hutcheon et al., 2010), we were able to determine that all of the FAD2 genes in each of the subgenomes were efficiently targeted by the Cas9/sgRNA complex.

To assess whether off-target mutations occurred, three sites in the genome most similar to the target FAD2 site were PCR-amplified and sequenced by Sanger chemistry using C. sativa DNA from C. sativa plants with a high mutation rate at the FAD2R1 target site. These three best off-target candidate loci contain a PAM and complete homology to the 10-bp ‘seed region’ that is at the 3’ end of the FAD2 protospacer (Jiang et al., 2015). In the 5’ ends, they contain five or six mismatches to the FAD2 protospacer (Figure S6). The resulting DNA sequence chromatograms from the Cas9-positive plants were indistinguishable from the WT chromatograms, suggesting that off-target mutations are not prevalent when using the FAD2 protospacer of interest (Figure S6).

Examination of the data in Data Set S2 also revealed that approximately 8% of the DNA sequencing reads contained a combination of single-nucleotide polymorphisms found in two (or, in one case, three) of the homoeologous FAD2 genes. Such
combinations could be due to unexpected recombination of homoeologous chromosome fragments following Cas9/sgRNA-generated chromosome breaks or, alternatively, to generation of incomplete DNA transcripts during polymerase chain reaction (PCR) DNA amplification and the subsequent utilization of these incomplete strands as templates during ensuing rounds of amplification. To distinguish between these two possibilities, the same DNA samples used for PCR amplification and Sanger DNA sequencing shown above were subjected to PCR amplification and DNA sequencing using Illumina Amplicon-Seq techniques in which thousands of DNA strands were sequenced from DNA isolated from each plant under study.

**PCR artefacts are generated during amplification of the three highly homologous copies of the FAD2 gene in Camelina**

The allohexaploid Camelina contains three distinct yet closely related homoeologous subgenomes, called A, B and C. Two subgenomes consist of seven chromosomes and the third of six chromosomes adding to the somatic chromosome content of 20 pairs (Hutcheon et al., 2010). Individual single-copy genes, such as the FAD2 gene, thus come in three allelic pairs that can be distinguished based on single-nucleotide polymorphisms (SNPs) specific to each subgenome (Hutcheon et al., 2010; Kang et al., 2011; Nguyen et al., 2013) (Data Set S2 and Figures 2 and 3). Our Sanger sequence data (Data Set S2) indicated that approximately 8% of the DNA sequencing reads contained a combination of single-nucleotide polymorphisms found in two (or, in one case, three) of the homoeologous FAD2 genes. Specifically, of the 259 Cas9/sgRNA-mediated mutations observed in our Sanger sequencing of cloned FAD2 gene DNA, 21 contained apparent chimeras between homoeologous chromosomes (Data Set S2). All possible combinations of homoeologous chromosome exchanges (A + C, B + C, etc.) were observed. No chimeric sequences were observed in the limited number of DNA sequences (i.e. 30) obtained from Sanger DNA sequencing of wild-type Camelina DNA samples.

**Figure 2** Clustal alignment of the sequences of the CsFAD2R1 target site in Camelina sativa leaf wild-type sample. Illumina Amplicon-Seq was used for DNA sequencing. Read frequency counts for each sample are provided to the left of the sequences. Only the sequences found at least 50 times are depicted. The three most common sequences represent the three WT homoeologs sequences. Asterisks denote nucleotides conserved in all three subgenome homoeologs (A–C). Lack of an asterisk denotes a nucleotide position at which there is a single-nucleotide polymorphism in one of the three homoeologs.
Because all homoeologs were targeted by the same guide RNA, it is possible that these sequences originated from recombination between homoeologous chromosomes following Cas9/sgRNA-generated chromosome breaks. Alternatively, as all sequences were amplified using the same primers, these sequences could have originated from chimeric PCR amplicons. To distinguish between these two possibilities, 30 transgenic plants expressing the Cas9 and sgRNA and two wild-type plants were selected from the samples analysed above and subjected to PCR amplification and DNA sequencing using Illumina Amplicon-Seq, generating thousands of sequences from each sample (Data Set S3 and S4). Analysed sequences were 350 bp long and amplified using nonhomoeologous specific primers, that is primers that perfectly matched all three homoeologous sequences. Paired-ended reads were obtained and processed using custom scripts (see Experimental procedures for details) to assign sequences to specific homoeologs, based on the SNPs distinguishing the three WT sequences. Chimeric sequences were observed in both transgenic and wild-type data sets (Data Set S3), and the percentage of chimeric sequence was at least as high in the control as in the transgenic samples (Table S2—summarized in Figure S3). These results thus provide no evidence for recombination between homoeologous DNA strands following Cas9/sgRNA-generated cleavage of the DNAs, but, rather, point to artificial recombination between homoeologous FAD2 gene strands during PCR amplification as the source of the observed chimeras. Reports in the recent literature (e.g. Liu et al., 2014 and references therein) document that PCR amplification of DNA from distinctly different, but highly homologous, DNA molecules can lead to the generation of ‘recombinant’ DNA strands due to a small number of incomplete PCR amplicons being used as primer sequences. These studies coupled with the present study point to the need for extreme care in interpreting data from experiments using PCR amplification of DNA from polyploid species.
CRISPR/Cas9 action results in a variety of mutant alleles

Sequencing the targeted sites allowed for an in-depth analysis of the patterns of mutations as, in total, 329 Sanger sequences and 410 490 Illumina sequences were analysed. Analyses of the initial Sanger data and the Illumina Amplicon-Seq data provided useful comparisons. First, the Amplicon-Seq data confirmed the presence of many different types of insertions and deletions at the CSFAD2R1 target site (average of 71%, indels) and the lack of insertions and deletions at the same site in DNA samples obtained from wild-type plant tissues (average 0.6% indels) (Figures 3 and 4 and Data Set S3). Next, it confirmed that all three homoelogous were efficiently targeted although the percentage of mutant sequence was higher in homoelogous B sequences (84.4%) than in homoelogous A (60.7%) or C (67.3%). Averaged across all mutated samples and homoelogous, Sanger reads indicated 21.2% WT, 78.2% indel-containing and 0.6% SNP-containing sequencing. These numbers were similar to those obtained from the Illumina reads, which indicated 27.5%, 71.1% and 1.4% of WT, indel-containing and SNP-containing reads, respectively. There were nearly twice as many Illumina reads containing insertions as compared to deletions. Within the Illumina reads, >99% of the insertions were single-nucleotide insertions with the following distribution: 49.1% T, 35.3% A, 12.9% G and 2.8% C. This distribution was consistent with observation from the Sanger data (50% T, 33% A, 12% G and 6% C), demonstrating strong insertion preferences. Deletions were more variable in both data sets. Of the 64 368 Illumina reads that contained deletions, half were 1-bp deletions, a quarter were 2-bp deletions, 11% were 3-bp deletions, and the remaining 12% were >3-bp insertions. For WT1, the Illumina reads were 92.2% WT, 0% indel-containing and 7.1% SNP-containing, and for WT2, they were 91.9%, 1.3% and 6.8%, respectively. The SNPs and indels observed in WT1 and WT2 are assumed to originate primarily from PCR-derived artefacts documented above and provide a base estimate of false positives.

Significant Cas9/sgRNA-mediated changes in seed oil composition

Analyses of fatty acid composition of seeds from both Arabidopsis and Camelina plants containing Cas9/sgRNA-mediated mutations in their FAD2 genes as well as from control plants lacking such modifications demonstrated a range of oleic acid content from the usual ~16% to greater than 50% (Data Set S5). One of the best-performing Arabidopsis lines (FAD2R2 T1-25-4 and its T4 seed progeny) showed an oleic oil concentration of ~50% in the T3 generation. This increased to a range between ~55% and ~60% in the T4 generation (Figure 4). As expected from results of earlier studies with Cas9/sgRNA-disrupted genes of Arabidopsis (Feng et al., 2014; Jiang et al., 2014), DNA sequencing of the FAD2 genes in T3 and T4 plants producing these high oleic acid seeds showed them to be homozygous for disruptions of the FAD2 genes (Figure S4). Also, as expected, the number of generations needed for oleic acid seed content to increase in the allohexaploid, Camelina, was more than in diploid Arabidopsis—although a few T4 Camelina seeds contained oleic acid levels of 50% or greater (Data Set S5 and Figure S5a). In each case in which there was a rise in oleic acid content, there was the expected proportional decrease in linoleic and linolenic acid content, leading to seeds with fatty acid composition more favourable for production of certain specialty oils and biodiesel. Targeting each of the three selected sites within the FAD2 gene (F1, R1 and R2) for Cas9/sgRNA attack was effective in causing significant increases in production of oleic acid and decreases in linoleic acid and linolenic acid (Figure S5a) in the best-performing Camelina lines. Production of total monounsaturated fatty acids (18:1, 20:1, 22:1) increased from ~30% in wild-type seeds to as high as ~74% (Figure S5b). These results demonstrate that, as with the use of RNAi, gene silencing and other gene editing technologies, Cas9/sgRNA system can be used to cause significant and favourable changes in seed fatty acid composition in both Arabidopsis and the emerging oil seed crop, Camelina. Consistent with our molecular off-target analysis, no evidence of overt off-target effects was observed in the phenotypes of plants or seeds containing Cas9/sgRNA-mediated gene mutations.

Changes in seed oil composition originate from the combined contributions of Cas9/sgRNA-mediated mutations in somatic cells of Camelina seeds and germ-line mutations

While expression of the Cas9 and sgRNA genes is clearly responsible for the changes in seed oil composition documented in Data Set S3, the data presented in Data Set S2 and S4 raise questions with regard to the source of this phenotype and the probability of its stable inheritance—at least in the present set of Camelina plants that contain and appear to maintain an active set of Cas9 and sgRNA genes. Careful analysis of the DNA sequences in the mutant genes shown in Data Set S2–S4 suggest, as detailed below, that while some mutant FAD2 genes observed in the present study may be inherited from one generation to the next, the large number of different mutations observed in DNA from a single sample of leaves or seeds from an individual plant points to a contribution of somatic cell mutations to the pool of DNA sequence variants observed. Specifically, cataloguing all types of mutations observed from the Amplicon-Seq data resulted in, on average, 59 different mutation types per sample, evenly distributed between the three homoelogous sequences (Data Set S4). We have also noted that the proportion of indel sequences was consistently higher in leaf samples (71.6%) compared to seed samples (58.5%), suggesting CRISPR/Cas9 activity is higher in somatic tissues than in germinal tissues (Data Set S3 and summarized in Figure S7). This is in agreement with our earlier observations and conclusions regarding Cas9/sgRNA-derived genetic mosaicism in somatic cells (Jiang et al., 2014) and similar
conclusions by others (e.g. Feng et al., 2014). Thus, the changes in fatty acid composition observed in individual seeds (Data Set S5) could be due to the effects of germ-line mutations combined with FAD2 mutations occurring in somatic cells during seed development. The latter would give rise to a mosaic of overlapping patches of somatic cells with varying combinations of mutations in the six copies of the FAD2 genes. This conclusion is further strengthened by our observations that when fatty acid composition is determined for individual T3 seeds from a single T2 plant or individual T4 seeds from a single T3 plant (Figures 4, 5 and Data Set S5), there can be noticeable variability in oleic acid levels from seed to seed (e.g. 39%, 40%, 43%, 45% and 55% for seeds from plant T1-5-1-1)—a potential reflection of a mosaic of somatic cells in individual seeds that carry variable numbers of inactivated FAD2 genes.

Germ-line mutations may be detected by the Mendelian inheritance of unique mutations in Cas9-positive plants or by the inheritance of Cas9-induced mutation in Cas9-negative plants. In the Amplicon-Seq data set from leaves of 27 individual Cas9-positive C. sativa plants, there are many cases in which a unique Cas9-induced mutation occurs in over 25% of the reads from a homologous pair (Data Set S4), suggesting that the mutation may have been inherited from one of the parents. On the other hand, many of these mutations are of a very common type and the high percentage could result from multiple somatic mutations as well. To better quantify the presence of germ-line mutations, we analysed Cas9-negative plants that were derived from a Cas9-positive C. sativa plant, T1-5 (Figure S8). Lack of the Cas9 gene in 20 nonfluorescent (i.e. ‘black’) seed progeny in each of three different T1-5 lines (i.e. T1-5-1-7B, T1-5-1-8B, T1-5-1-9B) was verified by PCR analyses (Figure S9). Sanger DNA sequence chromatograms indicate that some progeny from T1-5-1-8B contain a mutation in the B homoeolog and some progeny from T1-5-1-9B contain a mutation in the A homoeolog (Figure S10b). No mutation was detected in the other pools or in the C homoeolog (Figure S10c). From the pools that contained mutations, individual plants were sequenced and could easily be classified as ‘WT’, ‘homozygous A insertion’ or ‘heterozygous’ (Figure S10a). From these allelic ratios, we inferred that two of the 18 FAD2 homoeologous copies, or 11.1%, carried a mutation in the three Cas9-negative parents. Additionally, as demonstrated by this germ-line inheritance, we have generated Cas9-negative C. sativa plants that contain homozygous FAD2 knockouts in homologous pairs (Figure S10b).

While the presence of FAD2 gene knockouts in the germ-line may explain a substantial portion of the increase in oleic acid content observed, the high abundance of somatic cell mutations in developing seeds [documented by the large number of different Cas9/sgRNA-generated mutations in every line tested (Data Set S4)] suggests that FAD2 knockout mutations in somatic cells also may contribute to increases in oleic acid concentrations.

Cas9/sgRNA activity may contribute to changes in oil composition via mechanisms other than reduced expression. This is consistent with previous observations that FAD2 functions as a homodimeric enzyme (Lou et al., 2014) and that expression of nonfunctional mutants of FAD2 in cotton (Chapman et al., 2001, 2008) or closely related desaturases, epoxygenases, and hydroxylases in a variety of other plant species (Broun and Somerville, 1997; Broun et al., 1998; Cahoon et al., 1999; Singh et al., 2001) can cause inhibition of FAD2 desaturase activity—presumably

Figure 5 Oil profile in seeds of Camelina plants transformed using Cas9/sgRNA targeting the R1 site of the FAD2 genes. Variation of T3 seed oil profiles of individual seeds from T3 plant T1-1-3 (a) and T3 plant T1-5-1 (b); variation of seed oil profiles of individual T4 seeds from T3 plant T1-3-2-3 (c) and T3 plant T1-5-1-1 (d). Blue: wild-type control; light red: individual seed tested. Methods for extraction and measurement of Camelina seed fatty acids are detailed in Experimental procedures.
through formation of nonfunctional heterodimers (i.e., ‘subunit poisoning’). In these species, such dominant negative inhibition of FAD2 activity results in striking increases in oleic acid and decreases in linoleic and linolenic acids. (Broun and Somerville, 1997; Broun et al., 1998; Cahoon et al., 1999; Chapman et al., 2001, 2008; Lou et al., 2014; Singh et al., 2001). Thus, because the Cas9/sgRNA constructs used in the present studies primarily cause frame-shift mutations in the S’ region of the targeted FAD2 genes, it is likely that polypeptides are produced that contain authentic N-terminal FAD2 domains. It is possible that such molecules can binding with functional FAD2 monomers and partially inhibit desaturase activity. Future experimentation to test this hypothesis and, more importantly, future analyses of new generations of mutant Camelina plants containing homozygous or biallelic knockouts of FAD2 genes will be needed to provide a better understanding of the high oleic acid phenotypes presently observed.

A potentially significant advantage of using gene editing techniques such as ZFNs, TALNs and Cas9/sgRNAs is that once the desired gene editing event(s) have been obtained, the editing genes can be eliminated by simple cross-breeding techniques and the desirable phenotype stabilized by production of homozygous progeny (e.g. Li et al., 2012; reviewed in Weeks et al., 2015 and Petolino and Kumar, 2015). A lesson from the present study is that obtaining the desired phenotypic stability in some allohexaploid crops, such as Camelina, likely will require several generations in which the Cas9/sgRNA genes remain present and active. The rapid screening of seeds carrying active Cas9 and sgRNA genes by virtue of their easily detectable DsRed fluorescence is an attractive tool for achieving this goal. One potential future approach to gaining a better, but not complete, picture in regard to this uncertainty will be to analyse DNA from pollen samples from individual plants (or, if technically possible, from single pollen grains) to better determine the complement of wild-type and mutant FAD2 genes and the ratio between them. Alternatively, it is possible that germ-line mutations will be achieved more efficiently if the editing genes are driven by a different promoter, with documented expression in meristematic tissues. For example, homozygous mutants were recently obtained in a single generation in A. thaliana using a variety of promoters including an egg cell-specific promoter (Wang et al., 2015), the YAO promoter that is active in meristematic cells (Yan et al., 2015), the anther-specific DD45 gene promoter (Mao et al., 2016) and the INCURVATA2 gene promoter (Hyun et al., 2015).

Conclusions

Several important conclusions can be drawn from the present study. The Cas9/sgRNA system for gene editing is active in Camelina over multiple generations. Mutations in the FAD2 genes of Camelina can result in significant increases in oleic acid content and concomitant large decreases in linoleic and linolenic acid content. Based on results with other oil seed crops, future Camelina lines emanating from this continuing long-term project that are homozygous for inactivated FAD2 genes likely will exhibit even larger decreases in levels of long-chain polyunsaturated fatty acids and improved oleic acid content. Future projects aimed at producing plants that also contain Cas9/sgRNA-mediated knockouts of the fatty acid elongase, FAE1, genes should lead to additional boosts in oleic acid content of Camelina seeds.

Experimental procedures

Creation and analyses of transgenic Arabidopsis and Camelina plants and seeds: A. thaliana ecotype (Col-0) and C. sativa, cv. Suneson, were transformed using the floral dip method with vectors containing Cas9/sgRNA constructs targeting the respective Arabidopsis and Camelina FAD2 genes. Transgenic plants were self-pollinated, and leaves and seeds were collected from sequential generations of plants to determine by PCR/restriction enzyme analyses and DNA sequencing if gene editing events had occurred at the gene target sites. The fatty acid compositions of seeds were determined using gas chromatography. Mutant FAD2 gene DNA sequences (somatic and germ-line) in Camelina lines containing (Cas9-positive) and lacking (Cas9-negative) functional Cas9 and sgRNA genes were obtained by various combinations of Sanger DNA sequencing and Illumina Amplicon-MiSeq DNA sequencing techniques. A search for potential off-target gene modifications was conducted using Sanger DNA sequencing of three DNA sites most closely related to the target sites within the Camelina FAD2 genes. Details of each of these procedures are provided in Supporting Information.

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Author contributions

WJ developed methodology and performed or guided the experimentation. WJ and DPW designed the experiments. PL and IMH analysed and performed the Amplicon-Seq experiments. PL and IMH analysed the Amplicon-Seq data. WJ, DPW, EBC, LC and IMH interpreted the data and wrote the manuscript. DPW, EBC and LC are principal investigators on grants supporting this research.

References


transformed with a Brassica nonfunctional allele of a delta-12 fatty acid desaturase (FAD2). *Crops Sci.* **48**, 1470–1481.


Supporting information

Additional Supporting Information may be found online in the supporting information tab for this article:

**Figure S1** Enzymatic conversion of oleic acid to linoleic and linolenic acid by fatty acid desaturase 2, the enzyme whose gene, FAD2, is the target for inactivation by Cas9/sgRNA genes.

**Figure S2** PCR/Restriction Enzyme (PCR/RE) analysis of total DNA extracted from leaves of Arabidopsis T1 plants expressing Cas9/sgRNAs.

**Figure S3** Comparison of the rates of PCR recombinants during PCR amplification of Camelina FAD2 gene sequences using non-homologous specific amplification primers.

**Figure S4** Inheritance and homozygous condition of Cas9/sgRNA-mediated mutations at the R2 BbxCI target site of the

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FAD2 gene in T2 and T3 progeny of two transgenic Arabidopsis lines.

**Figure S5** Seed oil profiles in the best performing individual T4 seeds of Camelina plants transformed using Cas9/sgRNA targeting the R1, R2 and F1 sites in FAD2 genes.

**Figure S6** DNA sequencing of potential Off-Target sites.

**Figure S7** Model of CRISPR-Cas9 action in hexaploid Camelina.

**Figure S8** Lineage of Camelina sativa plants used to analyzed germline mutations.

**Figure S9** Detection of the Cas9 transgene sequence in 20 individual progeny (unlabeled lanes) of each of 3 black seeds.

**Figure S10** Detection of germline FAD2 mutations.

**Table S1** Primers upstream and downstream of the target sites in FAD2 genes in Arabidopsis and Camelina.

**Table S2** Rates of gene sequence chimera formation during PCR amplification of FAD2 gene DNA sequences.

**Appendix S1** DNA sequences of binary vectors used in this study.

**Data Set S1** Transgenic Arabidopsis and Camelina plants created for this study.

**Data Set S2** Overview of Sanger DNA sequencing data from Camelina FAD2 gene target sites.

**Data Set S3** Analyses of DNA sequences from wild-type Camelina plants and plants expressing CRISPR/Cas9.

**Data Set S4** Abundance of various mutations in wild-type Camelina plants and transgenic Camelina plants expressing CRISPR/Cas9.

**Data Set S5** Fatty acid composition of seeds from wild-type Arabidopsis and Camelina and from transgenic plants expressing CRISPR/Cas9.