A physical and genetic map of Cannabis sativa identifies extensive rearrangements at the THC/CBD acid synthase loci

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Cannabis sativa is widely cultivated for medicinal, food, industrial, and recreational use, but much remains unknown regarding its genetics, including the molecular determinants of cannabinoid content. Here, we describe a combined physical and genetic map derived from a cross between the drug-type strain Purple Kush and the hemp variety “Finola.” The map reveals that cannabinoid biosynthesis genes are generally unlinked but that aromatic prenyltransferase (AP), which produces the substrate for THCA and CBDA synthases (THCAS and CBDAS), is tightly linked to a known marker for total cannabinoid content. We further identify the gene encoding CBCA synthase (CBCAS) and characterize its catalytic activity, providing insight into how cannabinoid diversity arises in cannabis. THCAS and CBDAS (which determine the drug vs. hemp chemotype) are contained within large (>250 kb) retrotransposon-rich regions that are highly nonhomologous between drug- and hemp-type alleles and are furthermore embedded within ~40 Mb of minimally recombining repetitive DNA. The chromosome structures are similar to those in grains such as wheat, with recombination focused in gene-rich, repeat-depleted regions near chromosome ends. The physical and genetic map should facilitate further dissection of genetic and molecular mechanisms in this commercially and medically important plant.

[Supplemental material is available for this article.]
chromosome), and one or the other is inactivated in drug-type or hemp strains. This model was motivated by the discovery of a THCAS-like gene in hemp plants (Kojoma et al. 2006) and is consistent with the possibility that these related genes are derived from an ancient tandem duplication. In addition, physical linkage of genes involved in specialized metabolic pathways has been repeatedly observed in plants, similar to operons in bacterial genomes (Nützmann and Osbourn 2014); such a cluster was recently described for benzyloisouquinoline alkaloid biosynthesis genes in opium poppy (Guo et al. 2018). It is unknown whether genes involved in cannabinoid biosynthesis are clustered, although genetic analyses have previously indicated that at least one locus unlinked to THCAS/CBDAS contributes to cannabinoid content (Weiblen et al. 2015).

The draft genome and transcriptome of C. sativa described in 2011 (for a female plant of the drug-type strain Purple Kush [PK] and resequencing of a plant of the hemp variety “Finola” [FN]) (van Bakel et al. 2011) was unable to discriminate between these models due to high fragmentation. The C. sativa draft genome assembly, done largely with Illumina sequencing, was composed of 136,290 scaffolds, with an N50 of 16.2 kb. It was subsequently demonstrated that >70% of the C. sativa draft genome is composed of repetitive sequences (Pisupati et al. 2018). Measurement of single-nucleotide variants (SNVs) in four strains showed rates of heterozygosity ranging from 0.18%–0.26% and revealed that the drug-type and hemp-type strains were well separated by SNVs; the rate of occurrence of SNVs between these types was as high as 0.64% (van Bakel et al. 2011). Cytogenetic analysis has furthermore suggested a high degree of inter- and intraculivar karyotype polymorphisms (i.e., differences in homologous chromosomes that can be observed by microscopy), at least among hemp varieties (Razunova et al. 2016), which may further complicate genome assembly. To address these complications and to simultaneously leverage the high rate of SNVs between PK and FN, we coupled Pacific Biosciences (PacBio) long-read single-molecule real-time (SMRT) sequencing of PK and FN with Illumina resequencing of 99 F1 progeny between the two in order to generate a combined genetic and physical map. The combined map provides new insights into the arrangement of the chromosomes and the cannabinoid biosynthetic genes, including discovery of substantial rearrangement and gene duplications at the closely linked THC and CBD acid synthase gene loci.

### Results

A combined genetic and physical map reveals that genes and recombination events are concentrated near chromosome ends

We performed PacBio SMRT sequencing of genomic DNA (gDNA) from the female parent PK and the male parent FN to a depth of ~79× and ~98×, respectively. We used these data to develop an initial set of scaffolds, using the FALCON assembler (Chin et al. 2016), with PK and FN analyzed separately (Table 1). The assemblies were further polished with Illumina data using Pilon (Walker et al. 2014) to correct indel errors associated with homopolymer repeats in the PacBio data. The FN assembly was more contiguous than the PK assembly (scaffold N50 of 445.6 vs. 146 kbp, respectively), likely reflecting the increased FN coverage and the use of a more recent sequencing chemistry, and each substantially improved on our original Illumina assembly (Supplemental Fig. S1; van Bakel et al. 2011). De novo repeat classification using RepeatModeler (http://www.repeatmasker.org/RepeatModeler/) confirmed that the sequence of both assemblies is highly repetitive (~73%) (Supplemental Fig. S2), with hundreds of distinct families. The two sets of scaffolds largely mapped to each other one-to-one (Supplemental Fig. S3) but with differing breakpoints that mostly reflected differences in scaffold boundaries. The total size of the PK and FN assemblies was close to the haploid genome size estimated by flow cytometry (818 and 843 Mb for female and male, respectively) (Sakamoto et al. 1998). Overall, 90.3% of 30,074 previously

### Table 1. Genome assembly statistics

| Assembly statistics                          | PK                | FN                | FN anchored |
|---------------------------------------------|-------------------|-------------------|-------------|
| PacBio sequencing and assembly              |                   |                   |             |
| Total PacBio raw reads                      | 9,979,332         | 10,623,051        | N/A         |
| Total PacBio raw read bases (Gb)            | 64.62             | 82.32             | N/A         |
| Average PacBio raw read length (bp)         | 7179              | 8716              | N/A         |
| Total assembled bases (Mb)                  | 892               | 1009              | 784         |
| Scaffold N50 (kbp)                          | 146.0             | 445.6             | 382.9       |
| No. of scaffolds                             | 12,836            | 5303              | 2952        |
| Largest scaffold (Mb)                        | 1.41              | 2.49              | 2.49        |
| % PK transcriptome in genome (≥50% match)   | 90.3%             | 87.3%             | 78.5%       |
| % PK transcriptome in genome (complete)     | 82.7%             | 78.5%             | 70.4%       |
| % repeat content                            | 73.3%             | 73.9%             | 72.2%       |
| Haplotype blocks                             |                   |                   |             |
| FN haplotype blocks with >10 SNVs           | 34,197            | 13,098            | 10,557a     |
| No. of phased SNVs in haplotype blocks       | 2,734,893         | 1,359,019         | 1,214,845a  |
| % FN scaffolds with one or more haplotype block | 77.2%             | 86.7%             | 100%a       |
| % sequence in FN haplotype blocks           | 43.5%             | 76.1%             | 78.8%a      |
| FN haplotype block N50 (kbp)                | 27.6              | 92.6              | 98.7a       |
| No. of blocks used to create genetic map    | 14,440            | 4507              | N/A         |
| No. of SNVs used to create genetic map      | 1,888,187         | 799,227           | N/A         |
| Mean total coverage at SNVs used to create genetic map (parents and F1s) | 718 (+/− 350) | 660 (+/− 363) | N/A |
| Illumina sequencing                         |                   |                   |             |
| Total Illumina raw paired-end reads         | 105,000,000       | 162,968,810       | N/A         |
| Total filtered Illumina paired-end reads    | 80,369,366        | 98,244,687        | N/A         |
| Total filtered Illumina reads (Gb)          | 11.49             | 28.98             | N/A         |

*Statistics are based on the subset of FN haplotype blocks that are contained within scaffolds in the anchored map.
described PK transcripts (van Bakel et al. 2011) mapped to the PK assembly (82.3% mapping completely within a single scaffold). Each assembly also contained >95% of eudicotyledon single-copy orthologs from OrthoDB, of which >97% were complete (Supplemental Fig. S4), indicating that both assemblies represented the vast majority of the cannabis gene space. An ortholog duplication rate of >14% and slightly larger than expected assembly sizes suggest that some regions of the diploid genomes were resolved into separate contigs, which can be an issue for polymorphic species (Shimizu et al. 2017).

We reasoned that a genetic map would provide an independent means to link scaffolds, in addition to being independently useful for genetic analysis. To generate a genetic linkage map, we employed the SOILoCo pipeline, created by Scaglione et al. (2016) to create a map of the artichoke genome. We applied the pipeline to F1 data from a cross between a PK female and FN male. SOILoCo requires phasing of the parental scaffolds into blocks in which parental haplotypes can be uniquely identified. It then uses SNVs in the offspring to determine which of the parental haplotypes is inherited for each F1 at each block. The inherited parental haplotypes are called using a hidden Markov model, which compensates for uncertainty in genotype calling caused by relatively low coverage typical of resequencing, by taking advantage of the multiple SNVs in each block. Because each of the four parental haplotypes is traced uniquely, recombination frequencies between blocks (and thus between scaffolds) can subsequently be calculated, and the recombination frequencies can be used to place blocks (and scaffolds) into linkage groups. Since the blocks of informative SNVs differ between the parental types, a separate genetic map is created for each parent (in this case, PK and FN). In our implementation, we identified phased haplotype blocks of physically linked unique SNVs in the FN assembly contained within PK or FN PacBio raw reads using HapCUT2 (Table 1; Edge et al. 2017), and scored them in 99 F1 progeny using Illumina sequencing (median coverage about 4×). We then ran the SOILoCo pipeline, followed by R/qtl (Broman et al. 2003) and MSTmap (Wu et al. 2008), to form linkage groups and order scaffolds within them.

The blocks formed 10 large linkage groups in both PK and FN, which we assume correspond to the established nine autosomes and X/Y (which contain a pseudoautosomal region and recombine) (Peil et al. 2003) and are hereafter referred to as chromosomes. The maps were largely consistent between PK and FN (Supplemental Fig. S5) and were therefore merged (MergeMap) (Wu et al. 2011). The merged genetic map is depleted for short scaffolds, repetitive sequence, and scaffolds containing a higher proportion of SNVs with segregation distortion (these SNVs are ignored by SOILoCo). The merged map contains 2952/5304 scaffolds, 784/1006 Mb (78%) of the initial sequence, 89% of eudicotyledon single-copy orthologs, and 21,168/30,074 of all PK transcripts (70.4%) (Table 1; Supplemental Fig. S4).

Figure 1A plots composite physical versus genetic distance across the chromosomes, with several major trends in the chromosomal sequences also illustrated (Supplemental Fig. S5 shows similar graphs and also plots of genetic vs. physical distance, as well as a comparison of recombination frequencies, for all individual chromosomes). First, there is a very strong tendency for recombination to occur near chromosome ends, while there are typically large blocks lacking recombination events across the middle of the chromosome. Second, genes are much more frequent near chromosome ends. Because promoters and enhancers are typified by open chromatin, which appears to promote crossovers in diverse species, including maize (Liu et al. 2009) and Arabidopsis thaliana (Choi et al. 2013), this arrangement may underlie the observed recombination frequencies. Third, the poorly recombining central parts of chromosomes not only are gene-poor but also have a higher repeat content, which may be methylated and could suppress recombination (Zamudio et al. 2015). Fourth, assuming that the centromere is located within the nonrecombining central segments of the chromosomes, then Chromosomes 5, 9, and 10 appear to be telocentric (i.e., behave as if they have a single long arm). These may represent the sex chromosome, one end of which

![Figure 1](https://www.genome.org)

**Figure 1.** Comparison of physical and genetic distance in *Cannabis sativa* and arrangement of sequence features on chromosomes. (A) Median values are indicated for all metacentric linkage groups (Chromosomes 5, 9, and 10 are excluded), scaled to the same physical length. Black points indicate the median increase in genetic distance every 1/100th of the physical distance. Shaded histograms superimposed show density of repeat sequences. Density of genes and GC content are also indicated by blue and purple lines. (B) Values for Chromosome 6, which contains the THCAS/CBDAS loci; here, black points are the representative of individual scaffolds.
is nonhomologous and thus nonrecombining, and Chromosomes 8 and 9 (as determined by cytogenetics) (Divashuk et al. 2014), which harbor SS rDNA and 45S rDNA on one arm, respectively. The repetitive nature of these regions would be expected to impede both assembly and mapping. Indeed, four of five male-specific markers are found in the FN assembly, but none were placed on the genetic map, and the 45S and SS rDNA are not in the assembly (Supplemental Table S1).

Overall, the organization of *C. sativa* genes, repeats, and recombination frequency along chromosomes is similar to what is commonly observed in the grains (e.g., maize, barley, and wheat) (Gore et al. 2009; Liu et al. 2009; Mascher et al. 2017). To our knowledge, such an organization is unusual outside the grains: It has been observed in the walnut (Luo et al. 2015), but not thale cress (*A. thaliana*) (Meinke et al. 2009), apple (Di Pierro et al. 2016), strawberry (Davik et al. 2015), or mulberry (He et al. 2013), suggesting that this property is rare among Rosales.

### Genomic organization of cannabinoid pathway genes

We next examined the positions of genes encoding known cannabinoid biosynthetic enzymes on the chromosomes. With the exception of the functional copies of CBDAS and THCAS, which are considered below, the cannabinoid-related genes are distributed in a mostly random fashion across the genome (indicated in Supplemental Fig. S5). The new map also finds that *C. sativa* encodes one copy of AAEE (hexanoyl-CoA synthetase) and two tandem copies of tetraketide synthase ("olivetol synthase"). The genome sequences of both PK and FN also contain the THCAS-like gene described by Kojoma et al. (2006) which led to the two-locus THCAS/CBDAS hypothesis. This THCAS-like gene is 96% identical to THCAS at the nucleotide level and encodes a protein that is 93% identical to THCAS at the amino acid level. One copy of the THCAS-like gene is found in the PK assembly (scaffold 005500F: 2986–4620), and two are found in the FN assembly (scaffold 004887E: 13943–15377; 001793F, 69162–70796).

We examined the possibility that this THCAS-like gene encoded cannabichromenic acid (CBA) synthase (CBCAS), which is found in both drug-type and hemp strains and resembles THCAS and CBDAS in its catalytic mechanism (Morimoto et al. 1997). We expressed the predicted open reading frame as a secreted protein in *Pichia pastoris* strain X-33. We then added CBGA substrate to clarified culture media to test for enzyme activity. The products of this reaction were analyzed by high-performance liquid chromatography (HPLC), which revealed a specific signal for CBCA (Fig. 2A). Purification of the *Pichia* secreted protein through a series of chromatographic steps yielded a 59-kDa product at the expected size of CBCAS without its secretory signal sequence (calculated to be 58.9 kDa) (Fig. 2B). We next determined the kinetic properties of CBCAS after optimizing reaction conditions using the purified protein (Supplemental Fig. S6). At the optimal temperature of 40°C and a pH of 5.5, the reaction followed Michaelis–Menten reaction kinetics with a *K*<sub>m</sub> of 9.3 ± 2.3 µM and a *k*<sub>cat</sub> of 0.02 sec<sup>−1</sup>. These values are similar

![Figure 2](image-url) Characterization of CBCAS activity and expression. (A) HPLC analysis of CBCAS activity detected in *Pichia pastoris* cell cultures. Chromatograms of the CBGA substrate and CBCA standards are shown together with chromatograms of the enzyme reaction in media sampled from *Pichia* expressing CBCA in the presence of CBGA substrate before and after boiling at 95°C for 10 min. Insets correspond to the UV-absorbance spectrum (top) and the mass spectrum derived from a single quadrupole mass spectrometer (bottom) of the compound that eluted at 10 min. (B) SDS-PAGE analysis of CBCAS expressed in *P. pastoris* and purified by protein chromatography. (Lane 1) Protein ladder. (2) Concentrated protein fraction exhibiting CBCAS activity. The high-molecular-weight smear is glycosylated CBCAS. (3) Same fraction as lane 2, treated with EndoHf (MW = 70 kDa). (4) EndoHf only. (C) qRT-PCR analysis of CBCAS expression in cannabis tissues. cDNA derived from cannabis tissues was used as a template for PCR reactions using CBCAS-specific primers and EF1α as a reference gene. Differential expression of CBCAS is depicted as fold-change between tissue types compared with leaves. Trichome tissue consisted of isolated trichome secretory cells. (D) Quantification of CBCA content of the developing seedlings by HPLC.
to those reported for CBCAS purified from cannabis floral tissue ($K_m = 23 \mu M$, $k_{cat} = 0.04 \text{ sec}^{-1}$) (Morimoto et al. 1998). Finally, the accumulation of CBCA correlates well with the expression of CBCAS in various cannabis tissues, with the highest concentration observed in female floral tissue and minimal amounts in the leaf, stem, and root (Fig. 2C). Taken together, these data confirm that we identified the gene encoding CBCA synthase.

A previous study (Weibling et al. 2015) used QTL analysis in C. sativa to associate 121 genetic markers with total cannabinoid content and THCA/CBD ratio. Outside of THCAS/CBDAS, this study identified only one locus displaying a strong association with total cannabinoid content, at a distance of ∼1.2 cM between the trait and the marker. In our genetic map, this locus (marker ANUCS101) is linked to aromatic prenyltransferase (AP), which catalyzes the production of CBGA, the substrate of THCAS, CBDAS, and CBCAS, with a similar recombination frequency (2.1 cM in PK; 4 cM in FN). This observation suggests that either polymorphisms or differential regulation of AP contributes to cannabinoid production, presumably by controlling substrate concentration for THCAS and CBDAS. PK has greater than fivefold higher transcript levels of AP than FN (van Bakel et al. 2011), with no difference in copy number, suggesting that AP enzyme levels may be higher in drug-type plants partly due to differences in transcript levels. In addition to polymorphisms, there are multiple large (>100 bp) indels in and around the AP locus (including two within introns), which correspond mainly to LTRs, LINEs, and simple-repeat-like insertions, which could conceivably alter regulation of transcription or splicing (Fig. 3A).

Extensive rearrangement of the cannabinoid synthase locus underlies chemotype differences between PK and FN

Finally, we examined THCAS and CBDAS in the PK and FN genomes. The PK assembly contains only a single copy of THCAS and no exact copies of CBDAS: None have >95% identity to CBDAS at the nucleotide level. Similarly, the FN assembly contains only a single functional copy of CBDAS, while no THCAS gene is detected. These observations are confirmed by raw sequencing reads; no reads from FN map to THCAS, and no reads from PK map to CBDAS. Both genomes include the aforementioned CBCAS. This supports claims made using the draft genome and transcriptome (van Bakel et al. 2011). As expected from established segregation patterns, THCAS and CBDAS map to roughly the same region on Chromosome 6, near a known marker associated with THC and CBD content (ANUCS202) (Fig. 1B). However, the scaffolds that contain THCAS (in PK) and CBDAS (in FN) are dramatically different from each other, and neither has a clear counterpart in the other genome. The scaffold containing THCAS in PK does, however, contain a pseudogenic copy of CBDAS, with ∼94% identity to the known CBDAS sequence. The gene is likely nonfunctional as it has a gypsy element insertion at its center. Assuming these loci share common ancestry, there has clearly been extensive rearrangement since their divergence. The scaffold containing THCAS is ∼250 kb and that containing CBDAS is ∼750 kb, but the dotplot shown in Figure 3B illustrates almost complete lack of similarity over this span, with the exception of a large number of LTR-class retroelements. The extreme rearrangement clearly shows that these two genes do not have a simple isogenic relationship; Figure 3, A and C, illustrates more typical patterns of sequence similarity between PK and FN. The scaffold containing CBDAS is located within a much larger repeat-rich and gene-poor region of ∼49 Mb in the central section of Chromosome 6, encompassing 151 scaffolds with no recombination in either parent observed among the 99 F1s (Fig. 1B). The scaffold containing THCAS was separated from this region in a single recombination event among the 99 crosses, thus placing it at one end of this region and indicating that the THCAS and CBDAS scaffolds are at separate loci. We suggest that this repeat-rich segment of the chromosome may have hosted a series of tandem duplications and rearrangements amplifying an ancestral gene, leading to the present chromosomal organization; there is also a pseudogene with 89%–93% identity to each of THCAS, CBDAS, and CBCAS in this region. We note that this observation represents a modification of both previous models of CBDAS and THCAS arrangement: They are not isomorphic at an otherwise equivalent locus, and no equivalent of THCAS (deactivated or not) is found in hemp.

Discussion

The combined sequence/genetic map presented here is consistent with the known C. sativa karyotype and genome size, contains the vast majority of known transcripts, and largely correlates between PK and FN. To completely finish the sequence, it will most likely be necessary to further improve the resolution of the genetic map and/or leverage hybrid scaffolding technologies, e.g., by incorporating single-molecule genomic maps (Pendleton et al. 2015) or Hi-C data that provides >1 Mb phasing information (Kronenberg et al. 2018). Another future goal will be to identify and fully assemble the X/Y Chromosomes. There are numerous scaffolds in both PK and FN with no obvious counterpart in the other genome, which could represent distinctive components of the sex chromosomes and which were not captured in our genetic map.

The identification of CBCAS allows for a number of potential applications. Cannabichromene (CBC) is a weaker agonist of the cannabinoid CB1 and CB2 receptors compared with THC and CBD. However, unlike THC, both CBD and CBC have been shown to decrease nociception both by blocking the activity of ankyrin-type transient receptor potential channels that play roles in the perception of pain-inducing signals and by inhibiting the reuptake of endocannabinoids such as anandamide (Maione et al. 2011). Furthermore, CBC operates as a gastrointestinal anti-inflammatory agent in mice and protects adult neuronal stem progenitor cells in vitro (Izzo et al. 2012; Shinjyo and Di Marzo 2013). It therefore may be useful to breed medical cannabis strains with higher quantities of CBCA to treat specific ailments such as inflammatory bowel disease and Crohn’s disease. Finally, the high degree of sequence similarity between CBCAS, THCAS, and CBDAS and the presence of multiple pseudogenes suggest that gene duplication and divergence has been the key driver of cannabinoid end-product diversification in cannabis. Comparative sequence analysis of the enzymes will help ascertain which amino acids are important in catalysis, and may lead to the rational design of cannabinoid biosynthetic enzymes that produce novel cannabinoids not observed in nature.

Our identification of CBCAS also clarifies a puzzling finding of Kojoma et al. (2006), who used PCR to amplify a THCAS-like gene from “fiber-type” (hemp) cannabis that contained no THC. Based on the sequence of the gene that we show has CBCAS activity, the THCAS-like gene amplified by Kojoma et al. (2006) is CBCAS. This result makes sense, since nonnordrug/hemp forms of cannabis also contain CBCA.

Cannabis and cannabinoids are increasingly employed in medicine and recently have been legalized for recreational use in many jurisdictions. The new map should facilitate vastly improved
genetic analysis, including QTL mapping, which will accelerate crop improvement efforts. Drug prohibition has restricted access to cannabis by plant breeders and researchers, and as a result, it has received less attention than other crops. Cannabis suffers from insect pests and widespread fungal diseases and has a number of agronomic issues such as flowering time requirements that make it difficult to grow in some environments. In addition, breeding of cannabis types with specific cannabinoid and terpene profiles is desirable for the development of new varieties for medical and recreational use. The fact that a strong and interpretable result was obtained by re-examining a previously described marker correlating with total cannabinoid content (Weiblen et al. 2015) clearly shows the potential of this approach as it applies to cannabinoid metabolism. Due to the relatively high rate of polymorphism in cannabis, it should be possible to employ resequencing (e.g., low-coverage short-read Illumina protocols) either on crosses or at a population level to associate variants or variation with traits and genes, using the genetic map.

Methods

Plant cultivation and gDNA isolation

A female PK plant, produced through multiple vegetative propagation generations from the original source plant used to produce the draft *C. sativa* genome (van Bakel et al. 2011), was pollinated by a
male FN plant in an indoor growth chamber. Seeds produced from this cross were germinated under standard conditions and grown to seedling stage. gDNA was isolated from young leaves using a GenElute Genomic Miniprep Kit (Sigma-Aldrich). The secure facilities used for plant growing were licensed by Health Canada.

**PacBio SMRT sequencing of the PK and FN genomes**

gDNA library preparation and sequencing were performed according to the manufacturer’s instructions and reflect the P6-C4 sequencing enzyme and chemistry, respectively. PK and FN gDNA was first repurified using a 0.8× AMPure XP purification step (0.8× AMPure beads added by volume, to each DNA sample dissolved in 200 µL EB, vortexed for 10 min at 2000 rpm, followed by two washes with 70% alcohol and finally diluted in EB), to remove small fragments and/or biological contaminant. The purified DNA sample was taken through DNA damage and end-repair steps. Briefly, the DNA fragments were repaired using DNA damage repair solution (1× DNA damage repeat buffer, 1× NAD+, 1 mM ATP high, 0.2 mM dNTP, and 2× DNA damage repeat mix) with a volume of 21.1 µL and incubated at 37°C for 20 min. DNA ends were repaired next by adding 1× end repair mix to the solution, which was incubated at 25°C for 5 min, followed by the second 0.45× Ampure XP purification step. Next, 0.75 µL of blunt adapter was added to the DNA, followed by 1× template prep buffer, 0.05 mM ATP low, and 0.75 U/µL T4 ligase to ligate (final volume of 47.5 µL) the SMRTbell adapters to the DNA fragments. This solution was incubated at 25°C overnight, followed by a 65°C 10-min ligase denaturation step. After ligation, the library was treated with an exonuclease cocktail to remove unligated DNA fragments using a solution of 1.81 U/µL Exo III 18 and 0.18 U/µL Exo VII and then incubated at 37°C for 1 h. Two additional 0.8× Ampure XP purifications steps were performed to remove <1000-bp molecular-weight DNA and organic contaminant.

Size-selection was confirmed using the Agilent bioanalyzer, and the mass was quantified using a Qubit assay before proceeding with primer annealing and DNA sequencing. For PK, 100 pM of SMRTbell libraries were mag bead loaded and sequenced with a combination of P5/C3 and P6/C4 chemistry on a PacBio RSII machine with 6-hour movies. For FN, 3 pM of SMRTbell libraries were sequenced independently on the NextSeq 500 platform, yielding ≥49.9 Gbp.

**FALCON assembly and Illumina polishing**

FALCON (Chin et al. 2016) was used to generate genome assemblies for PK (v0.4.0) and FN (v1.8.6). Briefly, raw subread data were filtered to remove the shortest reads to an approximate coverage of 70× for each genome, leaving 8,003,220 (80.2%) of subreads for PK and 6,646,226 (62.6%) of subreads for FN, or ~58 Gbp for each. Paired-assembled reads (i.e., error-corrected reads) were then created with a length cutoff of ≥6000 bp for PK and ≥7000 bp for FN, resulting in 2,239,051 and 5,323,023 preassembled reads, respectively. The PK and FN genomes were then assembled using paired-assembled reads with a minimum length of 9 kbp or 7 kbp, respectively. Additional relevant assembly parameter settings for FN were as follows:

```
pa_HPCdaligner_option: -B128 -t16 -e0.8 -M24 -l1200 -k18 -h256 -w8 -s100 -T12
ova_HPCdaligner_option: -B128 -M24 -k24 -h600 -e.92 -l1800 -s100 -T12
falcon_sense_option: --output_multi --min_cov_aln 4 --min_idt 0.70 --min_cov 4
```

Similar assembly parameters were used for PK, except that min_cov was set to 3.

Each FALCON assembly was corrected with paired-end Illumina reads using Pilon version 1.22 (Walker et al. 2014) after mapping available Illumina sequencing data (van Bakel et al. 2011) to the FALCON-assembled genomes using BWA-MEM (version 0.7.8) (Li 2013) with an average of 96× (PK) and 23× (FN) coverage. Correction was performed with the “diploid” flag and the “bases” flag set to correct only indels and SNPs. A total of 1,511,828 insertions and 228,876 deletions were corrected in the FN assembly, and 1,807,453 insertions and 283,918 deletions were corrected in the PK assembly.

**Repeat content analysis**

Repeats in the FN and PK genomes were predicted de novo and classified using RepeatModeler (v1.0.11; http://www.repeatmasker.org/RepeatModeler/). RepeatModeler was applied to each assembly with the “nch” engine (RMBlast v2.2.28) provided with RepeatModeler. Other prerequisite components installed with the RepeatModeler package included RECON v1.0.8 and RepeatScout v1.0.5 (Price et al. 2005), Tandem Repeat Finder v4.0.4 (Benson 1999), and Repbase-derived RepeatMasker libraries (http://www.girinst.org/server/RepBase/) from January 2017. The de novo repeat classification provided by RepeatModeler was filtered to remove families with a >1-kb BLAT (Kent 2002) alignment to PK transcripts. The final filtered RepeatModeler output was then used as input for RepeatMasker (Smit et al. 2013–2015) to produce a masked version of the assembly and obtain the genomic positions of annotated repeats.

**Assessment of genome assembly completeness**

The completeness of each genome assembly was assessed using BUSCO v3.0 (Simão et al. 2015) and the set of eudicotyledons single-copy orthologs from OrthoDB v10, with default arguments in the provided virtual machine instance.

**Comparison of PK and FN scaffolds**

PK and FN assemblies were aligned using LASTZ (Harris 2007) version 1.04.00 with the -ungapped and -notransitions options and a step of 20. Alignments with an identity of ≤95% and a length of ≤2000 bp were removed. To produce a dotplot, FN contigs were initially ordered by size along the y-axis. Next, PK contigs were ordered and orientated on the x-axis by the position of their best hit on the y-axis. FN contigs were then reordered on the y-axis according to their best hit to the newly ordered contigs on the x-axis. This process was repeated until the order of contigs on the x-axis, and the order of contigs on the y-axis converged.

**Illumina sequencing of the FN and FI individuals**

Dual-indexed libraries were prepared using the Nextera DNA library preparation kit (Illumina), pooled equimolar, and sequenced on the HiSeq 2500 platform, yielding 529.9 Gbp total. FN was sequenced independently on the NextSeq 500 platform, yielding 49.9 Gbp.
Building the genetic map

**Quality filtering**
Barcode and adapter sequences were filtered from all FN and F1 Illumina PE reads. FN reads were further filtered using Sickle with the flags `-q 20 -l 125` (https://github.com/najoshi/sickle) (version 1.33). PK Illumina 2×100 PE reads from the 2011 draft genome were also filtered using Sickle, with the flags `-q 20 -l 90`.

**Variant calling**
BWA-MEM (Li 2013) was used to map Illumina paired-end reads for FN, PK, and the F1s to the PK FALCON assembly, after which Picard (http://broadinstitute.github.io/picard/) was used for sorting, duplicate marking, and indexing the alignments. To call variants for the F1s, we used the mpileup function from bcftools (Li et al. 2009) over all of the F1 individuals and both parents to overcome spots of lower coverage in the F1s. Variants were also called individually for each parent using the GATK HaplotypeCaller (McKenna et al. 2010) to be used as input for haplotype phasing.

**Phasing the parental haplotypes**
Haplotypes for the parents were phased using HapCUT2 (Edge et al. 2017), using the –pachio 1 argument to improve accuracy with PacBio reads and the –ea 1 argument to calculate switch quality scores. As input, parental SNPs called by HaplotypeCaller on the Illumina data were provided in conjunction with PacBio raw reads. This was done to both increase the length of the resulting haplotype blocks and boost confidence in the phasing by requiring agreement between the two sets of data. To further increase confidence, we only used SNPs that had a quality score greater than 25 and read coverage between six and 46 and that were more than five bases away from an indel. Haplotype blocks were then split if the switch quality score was less than 30. Finally, only blocks with more than 10 SNPs were retained to use as input for SOILoCo.

**Genotyping the F1s**
The SOILoCo method (Scaglione et al. 2016) was used to genotype the F1s at each haplotype block, using the output of HapCUT2 and the variants called by mpileup. Required values and divergence from the default parameters are as follows. For vcf2strings.pl, minor allele frequencies 1 and 2 (–MAF-1 and –MAF-2) were set to 0.25 and 0.75, respectively. This step allows the removal of any markers that may display segregation distortion (8.5% of markers show some degree of segregation distortion). As both scaffolds do not have a counterpart in the other parental assembly, genotypes were extracted from variant loci that meet the following criteria: an allele frequency of 0.5 in the parent harboring the scaffold, no coverage in the opposing parent, an allele frequency of 0.5 in the F1s, and all F1s are homozygous. The scaffold containing **THCAS** is the only scaffold from the PK FALCON assembly that was placed in the genetic map.

**Forming linkage groups**
R/qtl (Broman et al. 2003) was employed to divide haplotype blocks for each parent separately across linkage groups using the formLinkageGroups function with maximum recombination frequency (max.rf) set to 0.05 and minimum LOD (min.LOD) set to 15. The resulting linkage groups were compared against one another to identify any pairs of linkage groups with a mean recombination frequency of greater than 0.8 between the haplotype blocks they contain, in which case the switchAlleles function was used to swap the alleles for all the haplotype blocks in the smaller linkage group, and formLinkageGroups was called again. Afterward, R/qtl functions checkAlleles, switchAlleles, and formLinkageGroups were run in succession two more times to further identify and fix haplotype blocks with swapped alleles. All linkage groups with more than 100 haplotype blocks were passed to the ordering step. For PK, there were 11 linkage groups with more than 100 haplotype blocks; however, two of them just missed the cutoff for being joined together and were therefore combined. Further support for combining these linkage groups came from a comparison with the FN map, in which the scaffolds held in these two PK linkage groups were held in a single FN linkage group.

**Ordering scaffolds**
Haplotype blocks were ordered within each linkage group using MSTmap (Wu et al. 2008) with the Kosambi distance function. Three rounds of ordering were done with a smoothing step in between carried out using the Perl implementation of the SMOOTH correction algorithm (van Os et al. 2005) that is provided with the SOILoCo pipeline using an error threshold of 0.85. Correspondence between the two parental sets of linkage groups was determined based on similarity in the sets of scaffolds belonging to each linkage group. To handle ambiguity in scaffold placement, if the haplotype blocks for any given scaffold were distributed over more than one linkage group within or between parental maps, a census was taken to determine the correct linkage group, and haplotype blocks that did not agree with the majority were removed. If fewer than half of the haplotype blocks were in agreement, all haplotype blocks for that scaffold were removed, and the scaffold was not placed in either parental map. Finally, for each scaffold within each map, a distribution of the genetic positions (in cM) for all haplotype blocks belonging to the scaffold was established, and any outlier blocks were removed. After removal of ambiguous haplotype blocks and scaffolds, a final round of ordering was carried out for each parental map.

**Merging the genetic maps**
To translate each parental map from haplotype blocks to scaffolds in order that they could be merged, scaffold placements were determined by averaging the locations of the haplotype blocks belonging to each scaffold. The genetic maps for PK and FN were then merged using MergeMap (Wu et al. 2011) with the weight of the FN genetic map set to two and the weight of the PK genetic map set to one because it was based off the FN FALCON assembly.

**Gene cloning**
**CBDAS** was amplified from DNA isolated from FN leaves using gene-specific primers (forward: 5′-CTGAGGAAGACTATTGCATTGAGG-3′; reverse: 5′-AACGCTTCTATGTTTGGG-3′). PCR products were cloned.
into pCR8/GW/TOPO (Invitrogen), excised as PstI/Kpn1 fragments, and cloned into pPICz-alpa B (Invitrogen). The expression vectors were then transformed into P. pastoris strain X-33 (Invitrogen) by electroporation. Positive recombinants were selected for by plating transformed cells on YPD plates supplemented with 25 µg/mL phleomycin (Invivogen). To screen for activity, colonies were used to inoculate 5 mL BMG cultures, which were grown for 2 d at 37°C with shaking. The cells were then pelleted by centrifugation, resuspended in 5 mL BMM media, and grown for 4 d at 20°C with shaking with the addition of 1% methanol daily. Enzyme activity was tested by directly adding CBGA to clarified culture media, incubating overnight at 37°C, and then analyzing products by HPLC as previously described (Stout et al. 2012).

Quantitative PCR

RNA extraction, cDNA generation, and qRT-PCR conditions were identical to those previously reported (Stout et al. 2012). CBCAS primers (forward: 5′-CGGATGTACGTATGCTGCCA-3′; reverse: 5′-CATCTCTCATTAAATAGAAGAGCA-3′) were designed from alignments of THCAS-like genes identified in the cannabis genome to ensure their selectivity. Primers were tested using cloned THCAS, CBDAS, and CBCAS as templates. Any primer set that amplified a nontarget cDNA was discarded. Primer efficiencies were extrapolated from raw amplification data using LinRegPCR (Ruijter et al. 2009).

Recombinant CBCAS enzyme expression and purification

The culture with the highest CBCAS activity was selected for scaled up production. One milliliter of the initial culture was used to inoculate two 40 mL BMG cultures, which were grown for 2 d at 37°C. These cultures were then used to initiate two 400 mL modified BMM cultures that were buffered with 10 mM HEPES (pH 7) and were supplemented with rifobavin at 20 mg/L. These cultures were grown at 20°C with shaking at 100 RPM for 5 d, with methanol added to 1% by volume each day. The cultures were then clarified by centrifugation, and the resulting media were filtered and were passed over two Bio-scale Mini CHT hydroxyapatite cartridges (Bio-Rad) at a flow rate of 1.5 mL/min at 4°C. The cartridges were then attached in series to an AKTA FPLC system (GE Healthcare) and eluted with a 75-mL linear gradient of 5 mM sodium phosphate (pH 7) to 500 mM sodium phosphate (pH 7). Active fractions were pooled, concentrated with a 30 kDa cutoff Centricon filter (Millipore), and buffer exchanged into 20 mM citrate (pH 4.7) using a PD10 column (GE Healthcare). The resulting fraction was then injected onto a MonoS 5/50 cation exchange column (GE Healthcare) and eluted with a 75-mL linear gradient from 5 mM sodium phosphate to 500 mM sodium phosphate. Active fractions were pooled, concentrated with a 10 mL Amicon Ultra-15 filter (Millipore) and were supplemented with riboflavin at 20 mg/L. These cultures were then used to initiate two 400 mL modified BMM cultures that were buffered with 10 mM HEPES (pH 7) and were supplemented with rifobavin at 20 mg/L. These cultures were grown at 20°C with shaking at 100 RPM for 5 d, with methanol added to 1% by volume each day. The cultures were then clarified by centrifugation, and the resulting media were filtered and passed over two Bio-scale Mini CHT hydroxyapatite cartridges (Bio-Rad) at a flow rate of 1.5 mL/min at 4°C. The cartridges were then attached in series to an AKTA FPLC system (GE Healthcare) and eluted with a 75-mL linear gradient of 5 mM sodium phosphate (pH 7) to 500 mM sodium phosphate (pH 7). Active fractions were pooled, concentrated with a 30 kDa cutoff Centricon filter (Millipore), and buffer exchanged into 20 mM citrate (pH 4.7) using a PD10 column (GE Healthcare). The resulting fraction was then injected onto a MonoS 5/50 cation exchange column (GE Healthcare) and eluted with a 75-mL linear gradient from 5 mM sodium phosphate to 500 mM sodium phosphate. Active fractions were pooled, concentrated with a 10 mL Amicon Ultra-15 filter (Millipore) and were supplemented with riboflavin at 20 mg/L. These cultures were then used to initiate two 400 mL modified BMM cultures that were buffered with 10 mM HEPES (pH 7) and were supplemented with rifobavin at 20 mg/L. These cultures were grown at 20°C with shaking at 100 RPM for 5 d, with methanol added to 1% by volume each day. The cultures were then clarified by centrifugation, and the resulting media were filtered and passed over two Bio-scale Mini CHT hydroxyapatite cartridges (Bio-Rad) at a flow rate of 1.5 mL/min at 4°C. The cartridges were then attached in series to an AKTA FPLC system (GE Healthcare) and eluted with a 75-mL linear gradient of 5 mM sodium phosphate (pH 7) to 500 mM sodium phosphate (pH 7). Active fractions were pooled, concentrated with a 30 kDa cutoff Centricon filter (Millipore), and buffer exchanged into 20 mM citrate (pH 4.7) using a PD10 column (GE Healthcare). The resulting fraction was then injected onto a MonoS 5/50 cation exchange column (GE Healthcare) and eluted with a 40-mL linear gradient of 20 mM citrate (pH 4.7) + 500 mM NaCl. Active fractions were pooled, concentrated with a 30 kDa cutoff Centricon filter, and injected onto a Hiload 26/60 Superdex 200 size exclusion column (GE Healthcare). Proteins were eluted with a single column volume of 20 mM citrate (pH 5.0) + 150 mM NaCl. Throughout the purification, 1/10th volume of each fraction was retained for analysis to judge purity. Protein was isolated from each fraction using 15 µL of StrataCLean resin (Stratagene) and analyzed by SDS PAGE.

Enzyme assays and HPLC quantification of reaction products

To test for CBCAS enzyme activity during the protein purification, 150 µL of protein fraction was mixed with 50 µL of 500 µM sodium citrate buffer (pH 5.0) and 20 µmol of CBGA and incubated overnight at 37°C. The reactions were then extracted twice with ethyl acetate, and the organic fractions were pooled and dried in a SpeedVac concentrator. The products were then resuspended in 16 µL 50% methanol, of which 10 µL were analyzed by HPLC as previously described (Stout et al. 2012). Reactions for enzyme kinetic analyses were composed of 1 µg of purified CBCAS, 100 mM sodium citrate (pH 5.0), and 100 mM NaCl. These reactions were performed under Michaelis–Menten conditions at 40°C for 1 h. Reaction product extraction and analyses were the same as above.

Data access

The PacBio sequence read data generated for genome assembly, the Illumina sequencing data for the FN and F1 individuals, and the PK and FN genome assemblies from this study have been submitted to the NCBI BioProject database (http://www.ncbi.nlm.nih.gov/bioproject) under accession number PRJNA73819.

Competing interest statement

J.E.P. is the chief executive officer and shareholder of a for-profit cannabis science company, Anandia Laboratories, based in Vancouver, Canada. In this position he receives a salary. Anandia Laboratories performs analytical testing for licensed cannabis producers in Canada as well as works to develop new cannabis cultivars through breeding. L.H. was the chief research officer at CaniMed Therapeutics until May 2018. CanniMed is a for-profit company based in Saskatoon, Canada, that produces medical cannabis products for authorized patients. In this position he received compensation in the form of a salary and stock options. In June 2018 he moved to CB3 Life Sciences, where he is chief scientific officer. J.E.P. and J.M.S. have filed patent WO2015196275 on the nucleotide sequence encoding the enzyme CBCAS-based reagents, as well as methods for producing cannabinoids and/or altering cannabinoid production.

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References

Benson G. 1999. Tandem repeats finder: a program to analyze DNA sequences. Nucleic Acids Res 27: 573–580. doi:10.1093/nar/27.2.573
Broman KW, Wu H, Sen S, Churchill GA. 2003. R/qtl: QTL mapping in experimental crosses. Bioinformatics 19: 889–890. doi:10.1093/bioinformatics/19.6.889
Chin CS, Peluso P, Sedlacek HJ, Nattestad M, Concepcion GT, Clum A, Benson G. 1999. Tandem repeats finder: a program to analyze DNA sequence. Nucleic Acids Res 27: 573–580. doi:10.1093/nar/27.2.573
Dunn C, O’Malley R, Figueroa-Baldaras R, Morales-Cruz A, et al. 2016. Phased diploid genome assembly with single-molecule real-time sequencing. Nat Methods 13: 1050–1054. doi:10.1038/nmeth.4035
Choi K, Zhao X, Kelly KA, Venn O, Higgins JD, Yelina NE, Hardcastle TJ, Zielinski PA, Copenhaver GP, Franklin FC, et al. 2013. Arabidopsis meliotic crossover hot spots overlap with H2A.Z nucleosomes at gene promoters. Nat Genet 45: 1327–1336. doi:10.1038/ng.2766
van Bakel H, Stout JM, Cote AG, Tallon CM, Sharpe AG, Hughes TR, Page JE. 2011. The draft genome and transcriptome of Cannabis sativa. Genome Biol 12: R102. doi:10.1186/gb-2011-12-10-r102
van Os H, Stam P, Visser RG, van Eck HJ. 2005. SMOOTH: a statistical method for successful removal of genotyping errors from high-density genetic linkage data. Theor Appl Genet 112: 187–194. doi:10.1007/s00122-005-0124-y
Walker BJ, Abeel T, Shea T, Priest M, Abouelliel A, Sakthikumar S, Cuomo CA, Zeng Q, Wortman J, Young SK, et al. 2014. Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. PLoS One 9: e112963. doi:10.1371/journal.pone.0112963
Watt G, Karl T. 2017. In vivo evidence for therapeutic properties of cannabidiol (CBD) for Alzheimer’s disease. Front Pharmacol 8: 20. doi:10.3389/fphar.2017.00020
Weiblen GD, Wenger JP, Craft KJ, ElSohly MA, Mehmedic Z, Treiber EL, Marks MD. 2015. Gene duplication and divergence affecting drug content in Cannabis sativa. New Phytol 208: 1241–1250. doi:10.1111/nph.13562
Wu Y, Bhat PR, Close TJ, Lonardi S. 2008. Efficient and accurate construction of genetic linkage maps from the minimum spanning tree of a graph. PLoS Genet 4: e1000212. doi:10.1371/journal.pgen.1000212
Wu Y, Close TJ, Lonardi S. 2011. Accurate construction of consensus genetic maps via integer linear programming. IEEE/ACM Trans Comput Biol Bioinform 8: 361–394. doi:10.1109/TCBB.2010.35
Zamudio N, Barau J, Teissandier A, Walter M, Borsos M, Servant N, Bourchis D. 2015. DNA methylation restrains transposons from adopting a chromatin signature permissive for meiotic recombination. Genes Dev 29: 1256–1270. doi:10.1101/gad.257840.114

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