Developmental Plasticity of the Major Alkyl Cannabinoid Chemotypes in a Diverse Cannabis Genetic Resource Collection

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Cannabis is a chemically diverse domesticated plant genus which produces a unique class of biologically active secondary metabolites referred to as cannabinoids. The affinity and selectivity of cannabinoids to targets of the human endocannabinoid system depend on alkyl side chain length, and these structural-activity relationships can be utilized for the development of novel therapeutics. Accurate early screening of germplasm has the potential to accelerate selection of chemical phenotypes (chemotypes) for pharmacological exploitation. However, limited attempts have been made to characterize the plasticity of alkyl cannabinoid composition in different plant tissues and throughout development. A chemotypic diversity panel comprised of 99 individuals from 20 Cannabis populations sourced from the Ecofibre Global Germplasm Collection (ecofibre.com.au and anandahemp.com) was used to examine alkyl cannabinoid variation across vegetative, flowering and maturation stages. A wide range of di-/tri-cyclic as well as C₃-/C₅-alkyl cannabinoid composition was observed between plants. Chemotype at the vegetative and flowering stages was found to be predictive of chemotype at maturation, indicating a low level of plasticity in cannabinoid composition. Chemometric cluster analysis based on composition data from all three developmental stages categorized alkyl cannabinoid chemotypes into three classes. Our results suggest that more extensive chemical and genetic characterization of the Cannabis genepool could facilitate the metabolic engineering of alkyl cannabinoid chemotypes.

Keywords: Cannabis sativa L., hemp, medicinal Cannabis, LC-MS, propyl alkyl cannabinoids, tetrahydrocannabivarinic acid, cannabidivarinic acid

INTRODUCTION

Cannabis sativa L. is the sole, formally recognized species within the genus Cannabis and is a member of the angiosperm family Cannabaceae (Small and Cronquist, 1976). Cannabis is diploid (Van Bakel et al., 2011), predominately dioecious, and obligate outbred (Faeti et al., 1996) and can be considered highly heterozygote (Soler et al., 2017). The extant genepool is comprised principally of domesticated or previously domesticated feral populations...
The predominant bioactive secondary metabolites produced by Cannabis are the terpenophenolic phytocannabinoids (cannabinoids), of which >100 have been identified (ElSohly and Slade, 2005; Radwan et al., 2015). Structurally related terpenophenolic compounds also occur in other plant species such as the prenyllflavonoids in Humulus lupulus (Stevens et al., 1999), a closely related species within the Cannabaceae which is thought to have diverged ~21 MYA (Divashuk et al., 2014). However, the cannabinoids appear largely unique to Cannabis (Gertsch et al., 2010), and are formed at high concentrations within capitulate stalked trichomes on the floral tissues of female inflorescences. They also accumulate within capitulate-sessile trichomes and potentially bulbous trichomes on floral as well as non-floral tissues including leaves and stems (Happyana et al., 2013). Despite their relative abundance and interspersed distribution in plant tissue, the metabolic role of cannabinoids in Cannabis is largely unknown, although they may mitigate biotic stress via mitochondrial membrane dysfunction-induced necrosis in leaf cells (Morimoto et al., 2007).

Cannabinoids are produced in Cannabis in their carboxylic acid (COOH) forms and are decarboxylated to neutral cannabinoids in a non-enzymatic reaction which can be accelerated at temperatures >100°C (Dussy et al., 2005). Decarboxylation can also occur after extended periods of storage >100 days at room temperature (Hanuš et al., 2016). A notable example of this is the conversion of the non-psychoactive delta(9)-tetrahydrocannabinolic acid (THCA) to the psychoactive delta(9)-tetrahydrocannabinol (THC) (Izzo et al., 2009) upon loss of the COOH group.

The tricyclic THCA and dicyclic cannabidiolic acid (CBD) C₂₅-alkyl cannabinoids are the most predominant and commonly occurring cannabinoids in Cannabis (Figure 1; Hazekamp et al., 2016). A series of C₂₅-alkyl cannabinoid homologs, including the tricyclic delta(9)-tetrahydrocannabivarinic acid (THCVA) and dicyclic cannabidivarinic acid (CBDVA), can also contribute significantly to the cannabinoid profiles of ecotypes from Asian (Figure 1; Hillig and Mahlberg, 2004; Welling et al., 2016a) and African provenance (Baker et al., 1983), although these compounds are typically found at low levels in contemporary domesticated forms (Swift et al., 2013; Hazekamp et al., 2016; Welling et al., 2016a). Trace amounts of other alkyl homologs have also been identified such as methyl-(C₁) (Vree et al., 1972) and butyl-(C₄) (Smith, 1997) alkyl cannabinoids, although accounts of high levels of these cannabinoids in planta are scarce.

Current understanding of the bioactivity of cannabinoids is based on their modulation of the human endocannabinoid system, a poorly defined complex ensemble of several receptors, two endogenous cannabinoid ligands N-arachidonylethanolamine (anandamide) and 2-arachidonoylglycerol (2-AG) as well as associated enzymatic pathways (Di Marzo and Piscitelli, 2015). The cannabinoid alkyl side chain is a critical pharmacophore (Khanolkar et al., 2000), with changes in carbon length influencing the affinity and selectivity of plant derived cannabionoids to targets of the human endocannabinoid system (Thakur et al., 2005). Indeed, recent docking studies using a 2.6-Å resolution crystal structure of the human G-protein-coupled cannabinoid type-1 receptor (CB₁R) show binding of the tricyclic core of THC with a number of transmembrane domains preceding a highly conserved membrane-proximal N-terminal region, with the alkyl side chain extending toward a Trp356 κ.48 residual (Shao et al., 2016) associated with CB₁R activation (Shim et al., 2011). Subsequent partial agonist binding by THC to CB₁R stimulates mesolimbic dopamine activity (French, 1997), a mechanism believed to be partially responsible for this ligands psychoactivity.

Until recently, plant cannabinoids have primarily seen use in the context of recreational drug use of THC. However, they offer promise as novel therapeutics in a number of diverse non-communicable diseases. The company GW Pharmaceuticals, plc has developed cannabidiol (CBD) and THC containing Sativex (Chandra et al., 2017), a prescription medicine approved for the management of multiple sclerosis in more than 22 countries¹, as well as CBD containing Epidiolex² which has recently been approved by the US Food and Drug Administration (FDA) for the treatment of childhood seizures associated with Lennox-Gastaut syndrome and Dravet syndrome (Chandra et al., 2017). Ananda Hemp Ltd. (a subsidiary company of Ecofarm Industries Operations Pty Ltd.) has recently launched a range of cannabinoid-based products³. The C₃-alkyl cannabinoids cannabidiolvarin (CBDV) and delta(9)-tetrahydrocannabivarin (THCV) are also emerging as therapeutic entities. CBDV has been targeted by GW Pharmaceuticals, plc (Vemuri and Makriyannis, 2015), with phase I and II clinical trials having been initiated for the treatment of autism spectrum disorders and epilepsy, respectively. Moreover, a double-blind, placebo-controlled pilot study of 62 non-insulin treated type II diabetes subjects supports the therapeutic role for THCV in the modulation of fasting blood glucose and pancreatic β-cell function (Jadoon et al., 2016).

Current methods for the production of cannabinoid-based botanical drug products rely predominantly on clonal propagation of plants (Lata et al., 2012) due to the limited ability to predict chemical heritability in seed propagated progeny (Potter, 2014). Development of early diagnostic techniques to determine C₃-alkyl cannabinoid quality (CBDV + THCV) within the total cannabinoid fraction could assist breeders in the selection of elite alkyl cannabinoid breeding lines. While the ontogenetic variation in di-/tri-cyclic cannabinoid composition during plant development within the C₅-alkyl cannabinoid fraction has been studied (Pacifico et al., 2008; De Backer et al., 2012; Aizpurua-Olazïola et al., 2016; Richins et al., 2018), there have been limited attempts to characterize developmental changes of C₃-alkyl cannabinoid composition. Moreover, alkyl cannabinoid chemotypes have not been systematically evaluated among divergent subtaxa.

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This lack of clarity in understanding the extent to which alkyl cannabinoid composition varies in planta limits the ability to use chemotypic assessment during early developmental stages as well as to predict chemotype prior to seed formation. In the present study, liquid chromatography-mass spectrometry (LC-MS) profiling of a chemotypic diversity panel with a representative range of genotypes within the Cannabis genepool was used to characterize variation in alkyl cannabinoid composition across vegetative, flowering and maturation stages. Seed-based accessions were sourced from the Ecofibre Global Germplasm Collection with priority given to accessions with provenance from Southern, Eastern and Western Asia as well as Africa to ensure adequate representation of C$_3$-alkyl cannabinoid chemotypes (Hillig and Mahlberg, 2004; Welling et al., 2016a; Table 1). Cluster analysis of alkyl cannabinoid fractions was performed to provide insight into the categorization and genetic regulation of alkyl cannabinoid chemotypes in Cannabis.

**MATERIALS AND METHODS**

**Genetic Resources**

Acquisition, storage and experimental endeavors were performed under the provisions of the Drug Misuse and Trafficking Act 1985 and in accordance with authorizations granted by the New South Wales Ministry of Health, Pharmaceutical Regulatory Unit, Legal and Regulatory Services Branch, Australia. Seed accessions were obtained from the Ecofibre Global Germplasm Collection owned by the company Ecofibre Industries Operations Pty Ltd. and managed by Southern Cross University, Australia. A single seed pack accession in a Cannabis genetic resource base collection can be generated from multiple parents and so is provisionally considered as a population (Faeti et al., 1996). Twenty populations (accessions) with geographical origins associated with C$_3$-alkyl cannabinoid accumulation (Hillig and Mahlberg, 2004) were preferentially selected to ensure an adequate level of alkyl cannabinoid chemotypic diversity (Table 1).

**Growth Parameters**

Growth parameters followed those of Welling et al. (2016a). Seeds were planted at a depth of 1.5 cm in cells of 5 cm (diameter) × 6 cm (height) in a mix of one part vermiculite, one part perlite, peat moss, and dolomite (110g/100L). CANNA® Aqua Vega nutrient solution was used as a supplement. Seedling trays (40 cells) were watered with 500 mL of water three times per day for 14 days. Seedlings were transplanted to 8 L pots, with each pot containing 8 g of Micromax® micronutrient formula and 100 g Osmocote® Exact slow release nutrient mix. Plants were grown in chambers fitted with ‘smart valves’ to maintain optimal water regimes. Temperature was maintained between 26 and 28°C, and plants were subject to 11 h of high pressure sodium (HPS)/metal halide (MH) light (luminous flux = 72,000 lumens) per day.

A total of 99 individual female plants were chemotyped at three developmental stages, with three to seven plants analyzed per accession (Table 1). Developmental stages were determined from visual inspection of plant morphological changes defined...
were dried at 35°C in air sealed containers with 3–5 mm orange silica gel beads. 

Approximately 250 mg of dried leaf tissue was extracted in 25 mL of HPLC grade EtOH (100%) for 30 min. To remove particulate material, 1 mL of the extract was centrifuged using a Compact centrifuge 2–5 (Sigma) at 3000 rpm for 10 min. The supernatant (600 µL) was transferred into 2 mL screw cap glass vial and all samples were subject to a 1:5 dilution to ensure signals were within calibration range.

### LC-MS Cannabinoid Profiling

Liquid chromatography-mass spectrometry (LC-MS) cannabinoid profiling runs were conducted using an Agilent 1290 Infinity analytical HPLC instrument (Agilent Technologies, Palo Alto, CA, United States), comprising of a vacuum degasser, autoinjector, binary pump and diode array detector (DAD, 1260), coupled with an Agilent 6120 Single Quadrupole mass detector (MSD). The LC-MS instrument was controlled using Agilent ChemStation software (Rev. B.04.03 [54]). Absorbance was monitored at 210 nm, 214 nm, 272 nm, 280 nm, 330 nm and 360 nm. An Agilent Eclipse plus rapid resolution high definition (RRHD) C18 column (1.8 µm; 50 mm × 2.1 mm internal diameter) was used and column temperature was set at 30°C. Injection volume was 3 µL.

The mobile phase followed those of Giese et al. (2015) with minor modification. Mobile phases consisted of 0.005% TFA in Milli-Q® water for channel A and 0.005% TFA in acetonitrile for channel B. Flow rate was 0.3 mL/min starting with an isocratic phase at 66% B for 8 min, then a linear gradient to 95% B over 4 min. 95% B was held for 1 min, then re-equilibrated to 66% B for 1 min. Equilibration was further extended for 1 min to perform an internal needle wash of the autosampler to minimize carryover. Run time was 16 min.

### Sample Preparation and Extraction

Disruption of fresh leaf tissue was performed using a Qiagen TissueLyser®. Frozen leaf tissue was ground in a 2 mL Eppendorf® Safe-Lock microcentrifuge tube containing a 3 mm Qiagen Tungsten Carbide Bead (Cat No./ID: 69997). Microcentrifuge tubes were agitated at 30 rotations per sec for 2 × 30 s intervals. Tissue was extracted in 1 mL of high-performance liquid chromatography (HPLC) grade EtOH (100%). Extractions were vortexed and mixed by agitation for 30 min. To remove particulate material, samples were centrifuged using a Compact centrifuge 2–5 (Sigma 113) at 8000 rpm for 30 min. The supernatant (600 µL) was transferred into 2 mL screw cap glass vial and subject to a 1:5 dilution to ensure signals were within calibration range.

Sample preparations for dried leaf tissue followed those of De Backer et al. (2009) and Welling et al. (2016a) with slight modification. Dried leaf tissue was ground with a Mixer Mill MM 301 (Retsch GmbH) at 30 rotations per sec for 30 s intervals. Duplicate extracts were performed for each plant per accession.

| Accession | ID | Individuals (n) | Provenance | Taxon | Source |
|-----------|----|----------------|------------|-------|--------|
| EIO.MW15.A | A  | 3              | Southern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.B | B  | 4              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.C | C  | 5              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.D | D  | 3              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.E | E  | 5              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.F | F  | 5              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.G | G  | 6              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.I | I  | 6              | Southern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.J | J  | 6              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.K | K  | 4              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.L | L  | 4              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.M | M  | 7              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.O | O  | 6              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.P | P  | 6              | Eastern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.Q | Q  | 5              | Caribbean    | Cannabis sativa L. | EFGGC |
| EIO.MW15.R | R  | 6              | Southern Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.S | S  | 3              | Southern Africa | Cannabis sativa L. | EFGGC |
| EIO.MW15.T | T  | 5              | Western Asia | Cannabis sativa L. | EFGGC |
| EIO.MW15.U | U  | 7              | Eastern Africa | Cannabis sativa L. | EFGGC |
| EIO.MW15.X | X  | 3              | Eastern Asia | Cannabis sativa L. | EFGGC |

EFGGC, Ecofibre Global Germplasm Collection.

in the Decimal Code for Growth Stages of Hemp (Mediavilla et al., 1998). Two × 250 mg fresh plant material was collected from the sub-apical raceme of each individual at opposing phyllotaxis during vegetative (fourth leaf pair, code 1008) and alternate phyllotaxis during flowering (code 2202) stages. Fresh leaf material was snap-frozen using liquid nitrogen in 2 mL Eppendorf® Safe-Lock microcentrifuge tubes and stored at −80°C. At seed maturation (code 2202) individual plant racemes were dried at 35°C in a forced ventilation oven for 72 h and stored at room temperature in air sealed containers with 3–5 mm orange silico gel beads.
MSD parameters followed those of Liu et al. (2014) and Welling et al. (2016a) with modification to allow quantification of four additional cannabinoids; THCV A, CBDV A, CBDV and cannabichromene (CBC). The MSD was operated in atmospheric pressure electrospray ionization mode (AP-ESI); scan mass range, 100-1200; drying gas temperature, 350 °C; fragmentor, 150; capillary voltage, 3000 V (positive); vaporizer temperature, 350 °C; drying gas flow, 12 L/min (N2); nebulizer pressure, 35 psi.

Quantification of cannabinoids was performed using selected-ion monitoring (SIM) with four available MSD signal channels (Supplementary Table S1). THCA, THC, THCV, cannabidiol (CBN), cannabigerolic acid (CBGA), cannabigerol (CBG), CBDA, CBD, CBVD, and CBC cannabinoid standards were sourced from Novachem Pty Ltd. (Melbourne, VIC, Australia). THCV A and CBDV A were isolated from plant tissue to develop analytical standards. All cannabinoid reference standards were scanned in positive mode [M + H]+ to determine the most abundant and representative signal.

Quadratic regression of calibration curves of individual reference standards was used to determine cannabinoid concentrations. Calibration curves were obtained from six solutions comprising of five acid cannabinoid standards THCA, CBDA, CBGA, THCV A, and CBDA at the following concentrations; 0.032, 0.16, 0.8, 4, 20, and 100 µg/mL. Calibration curves were also obtained from six solutions comprising of seven neutral cannabinoid standards THC, THCV, CBN, CBG, CBD, CBVD, and CBC at the following concentrations; 0.032, 0.16, 0.8, 4, 20, and 100 µg/mL. Linear regression analysis showed calibration curves to be linear within the concentration range for each cannabinoid (R² > 0.99). To minimize MSD interday variability, calibration curves were performed daily. The precision of the MSD was examined by injecting standard solutions six times within a 24 h period and relative standard deviation (RSD) for each cannabinoid peak area was <2%.

Statistical Analysis
To test for repeatability between extraction replicates, the C3-alkyl (FCS), C5-alkyl (FC5), dicyclic (Facyclic), and tricyclic (Ftricyclic) cannabinoid fractions were calculated using R². Strong positive correlations between extraction replicates were found for the FCS/FCS values (R² > 0.99) as well as for the Facyclic/Facyclic values (R² > 0.99) at vegetative, flowering and maturation stages. As such, mean values gathered from duplicate extraction replicates were utilized for statistical analysis. Statistical analysis was performed using GenStat 64-bit Release 18.1 (VSN International Ltd.) software. For regression analysis, the constant (intercept) was omitted and the fitted line was constrained through the origin. For non-hierarchical k-means cluster analysis, similarities were calculated using Euclidean distance.

Isolation, Purification, and Structural Elucidation of C3-Alkyl Cannabinoids
Dried female Cannabis floral tissue (4 x 1 g) sourced from the Ecofibre Global Germplasm Collection was extracted in 100% MeOH (4 x 20 mL) and evaporated using a Christ® BETA-RVC rotational vacuum concentrator. Extracts were pooled, resuspended in MeOH (4 mL) and partitioned using n-hexane (4 mL) to remove chlorophyll. The MeOH fraction was separated using a glass pipette, centrifuged to remove particulate matter and evaporated using a Christ® BETA-RVC rotational vacuum concentrator. The crude MeOH fraction (486 mg) was then resuspended in 6:4 MeOH: Milli-Q® water (2 mL).

Isolation and purification of the crude Cannabis MeOH extract was performed using an Agilent 1260 Infinity preparative HPLC system, comprising of a vacuum degasser, autosampler, binary preparative pump, diode array detector (DAD, 1260) and analytical-scale fraction collector. The preparative HPLC instrument was controlled using Agilent ChemStation software (Rev. B.04.03 [16]). Absorbance was monitored at 210 nm, 254 nm, 272 nm, 280 nm and 360 nm. A Luna C18 column (5 µm; 150 mm x 21.20 mm internal diameter) was used. Injection volume was 500 µL. Mobile phases consisted of 0.05% TFA in Milli-Q® water for channel A and 0.05% TFA in acetonitrile for channel B. Flow rate was 20 mL/min, starting with an isocratic phase at 80% B for 3 min, then a linear gradient to 99% B over 5 min. 99% B was held for 5 min, then re-equilibrated to 80% B for 2 min and held at 80% B for 5 min. Run time was 20 min. The fraction collector was operated in time-based trigger mode at 0.18 min time slices. THCV A (1.57 mg) and CBDV A (1.83 mg) fractionations were evaporated using a Christ® BETA-RVC rotational vacuum concentrator and redissolved in HPLC grade EtOH (100%).

Structural elucidation of C3-alkyl cannabinoids THCV A and CBDV A was performed using a Bruker Avance III HDX 800 MHz spectrometer. LC-MS spectra were obtained using an Agilent 1290 Infinity analytical HPLC instrument (Agilent Technologies, Palo Alto, CA, United States), comprising of a vacuum degasser, autoinjector, binary pump and diode array detector (DAD, 1260), coupled with an Agilent 6120 Single Quadrupole MSD. UV spectra were monitored at 210, 272, 280 and 360. For two dimensional NMR, 1H-1H Correlation Spectroscopy (1H-1H-COSY), Heteronuclear Single Quantum Coherence (HSQC), Heteronuclear Multiple Bond Correlation (HMBC), and Rotating-Frame Overhauser Spectroscopy (ROESY) experiments were performed. Data analysis, acquisition and processing of NMR and LC-MS spectra was conducted using TopSpin™ (TS3.5pl6) and Agilent ChemStation® (Rev. B.04.03 [54]) software, respectively.

RESULTS
Structural Elucidation of Acidic C3-Alkyl Cannabinoids
At the time of analysis, analytical standards for THCV A and CBDV A were not commercially available. Unknown compounds 1 and 2 were isolated and purified from Cannabis floral tissue, with structural elucidation performed using LC-MS (Supplementary Figures S1, S2) as well as 1H (Supplementary Figures S3, S4) 13C NMR (Supplementary Figures S5, S6) and 2D NMR (Supplementary Figures S7–S14). AP-ESI MS spectra of 1 and 2 exhibited the expected molecular ion m/z 328.9
opened pyran ring configuration of C and confirmed that both compounds were acidic cannabinoids. The was characteristic of a COOH at C-2 of 1 as well as the absence of a -OH group at associated positions (Supplementary Figures S11, S12 and HMBC (Figure 2). The presence of signals $\delta_C$ 173.9 (2-COOH) (1) and $\delta_C$ 174.2 (2'-COOH) (2) (Supplementary Figures S5, S6) as well as the absence of a -OH group at associated positions was characteristic of a COOH at C-2 of 1 and C-2' of 2, which confirmed that both compounds were acidic cannabinoids. The ROESY spectrum suggested a trans relationship between H-6a and H-10a of 1 as well as H-4 and H-3 of 2 (Supplementary Figures S13, S14). Compounds 1 and 2 were subsequently defined as THCVA and CBDVA, respectively.

Distribution of the Major Cyclic and Alkyl Cannabinoid Chemotypes

Chemotypes of 99 individual Cannabis plants from 20 seed accessions were characterized across three developmental stages using LC-MS analysis. Fresh leaf tissue samples were taken at the vegetative and flowering stages and cannabinoid composition was compared with dried floral tissue cannabinoid composition at maturation. The dicyclic cannabinoids cannabichromenic acid (CBCA) and cannabichromevarinic acid (CBCVA) as well as the precursor C3-alkyl cannabinoid cannabigerovarinic acid (CBGVA) were not commercially available at the time of analysis, nor were these compounds present at sufficient quantities to develop analytical standards. THCA, CBDA, THCV, and CBDVA as well as corresponding neutral decarboxylated derivatives were used as a proxy for C3-alkyl (FC3) and C5-alkyl (FC5) as well as dicyclic (FDicyclic) and tricyclic (FTricyclic) cannabinoid fractions within the total cannabinoid fraction. Calculation of the total cannabinoid fraction was achieved by the addition of THCA, CBDA, THCV, and CBDVA as well as their neutral cannabinoids (Supplementary Table S2). To determine the total cannabinoid fraction and to compare the FC3, FC5, FDicyclic, and FTricyclic values between juvenile and mature plants, neutral cannabinoids CBDV, CBD, THCV, and THC were expressed as acidic cannabinoids using formulae which accounted for differences in molecular weight:

$$F_{C3} = \frac{((\text{THCVA} + \text{CBDVA}) + ((\text{THCV} + \text{CBDV}) \times 1.1536))}{\text{total}} \times 100$$

$$F_{C5} = \frac{((\text{THCA} + \text{CBDA}) + ((\text{THC} + \text{CBD}) \times 1.1399))}{\text{total}} \times 100$$

$$F_{\text{dicyclic}} = \frac{((\text{CBDVA} + \text{CBDDA}) + ((\text{CBDV} \times 1.1536) + (\text{CBD} \times 1.1399)))}{\text{total}} \times 100$$

$$F_{\text{tricyclic}} = \frac{((\text{THCVA} + \text{THCA}) + ((\text{THCV} \times 1.1536) + (\text{THC} \times 1.1399)))}{\text{total}} \times 100$$

At maturation, variation in chemotype appeared to segregate within the accessions and so chemotype was reported at the plant level (Figure 3A), although within-accession chemotypic variation was more evident from the FDicyclic values than from the FC3 values (Figure 3A). Distributions of the di-/tri-cyclic as well as the C3-/C5-alkyl cannabinoid fractions at maturation were skewed toward high FTricyclic and FC5 values, respectively (Figure 3B). A wide range of the di-/tri-cyclic as well as the C3-/C5-alkyl cannabinoid fractions was found within the chemotypic
diversity panel derived from the Ecofibre Global Germplasm Collection, with Fc values ranging from 0.43% (±0.00%) to 87.78% (±0.10%) (Figure 3B). Plants from the Ecofibre accessions E as well as P (Ecofibre proprietary line) had the highest proportions of dicyclic (CBDVA) and tricyclic (THCVA) C3-alkyl cannabinoids, respectively. The plant from accession E with the highest dicyclic C3-alkyl cannabinoid fraction exhibited 81.2% CBDVA (% total cannabinoids), while the plant from accession P (Ecofibre proprietary line) with the highest tricyclic C3-alkyl cannabinoid fraction exhibited 75.1% THCVA (% total cannabinoids). Three discrete distributions comprised of low Fdicyclic/FTricyclic, intermediate Fdicyclic/FTricyclic, and high Fdicyclic/FTricyclic ratios were observed (Figure 3C), while the C3-/C5-alkyl cannabinoid proportions/ratios presented as a continuum with no obvious distribution patterns (Figures 3B,C).

**Stability of Alkyl Cannabinoid Composition**

A simple linear regression model was calculated to predict the di-/tri-cyclic as well as the C3-/C5-alkyl cannabinoid fractions at maturation based on cannabinoid fractions at vegetative and flowering stages. Regressions were significant at the vegetative stage for the Fdicyclic values [F(1, 98) = 15772.31, \( p < 0.001 \)], with an \( R^2 > 0.991 \), as well as for the FC3 values [F(1, 98) = 4301.82, \( p < 0.001 \)], with an \( R^2 > 0.964 \) (Figure 4A). Cannabinoid fractions showed minimal plasticity throughout development, with significant regressions also found at the flowering stage for the Fdicyclic values [F(1, 98) = 50480.89, \( p < 0.001 \)], with an \( R^2 > 0.997 \), as well as for the FC3 values [F(1, 98) = 8488.54, \( p < 0.001 \)], with an \( R^2 > 0.982 \) (Figure 4A).

As the di-/tri-cyclic as well as the C3-/C5-alkyl cannabinoid fractions approached parity in the vegetative stage, they appeared less predictive of chemotype at maturation when compared with cannabinoid fractions at the flowering stage (Figure 4A). To examine this further we truncated the Fdicyclic (n = 20) as well as the FC3 (n = 41) values by removing chemotypes with cannabinoid values of >90%/<10% and performed stepwise deletion of the data points with the largest standardized residuals (Figure 4B). For the di-/tri-cyclic cannabinoid fractions, three plants M01, M02, and M04 from the East Asian accession M contributed to reducing the explained variance between vegetative and maturation stages by 42.0% (Figures 4B,C), whereas for the C3-/C5-alkyl cannabinoid fractions, the removal
FIGURE 4 | (A) Regression analysis of the $F_{\text{dicyclic}}$ as well as the $F_{C3}$ chemotypic values between developmental stages. $F_{\text{dicyclic}}$ as well as the $F_{C3}$ values describe dicyclic as well as C3-alkyl cannabinoid fractions. The $F_{\text{dicyclic}}$ as well as the $F_{C3}$ chemotypic values on the x-axis describe cannabinoid fractions at the vegetative and flowering stages. (B) Regression analysis of the truncated $F_{\text{dicyclic}}$ as well as $F_{C3}$ values between developmental stages. The $F_{\text{dicyclic}}$ as well as the $F_{C3}$ chemotypic values on the x-axis describe dicyclic as well as C3-alkyl cannabinoid fractions at the vegetative and flowering stages. (C) Individual plants with large standardized residuals across vegetative and maturation growth stages. The $F_{\text{dicyclic}}$ as well as the $F_{C3}$ chemotypic values on the y-axis describe dicyclic as well as C3-alkyl cannabinoid fractions across developmental stages. Red arrow indicates position of units with large standardized residuals; Letters specify accession ID (Table 1); Numbers indicate plant individual within accession; $C_5$-alkyl cannabinoid fractions ($F_{C5}$); $C_3$-alkyl cannabinoid fractions ($F_{C3}$); dicyclic cannabinoid fractions ($F_{\text{dicyclic}}$); and tricyclic cannabinoid fractions ($F_{\text{tricyclic}}$).
Chemometric Categorization of Alkyl Cannabinoid Composition

Chemometric categorization of the di-/tri-cyclic as well as the C₃-/C₅-alkyl cannabinoid fractions was performed using non-hierarchical k-means cluster analysis which incorporated within-plant variation across vegetative, flowering and maturation developmental stages. This was based on the premise that the genotype does not vary over time, and that the continuity of the C₃-/C₅-alkyl cannabinoid fractions could be disentangled by removing non-genotypic contributions to chemotype. The optimal number of clusters based on criterion values as a function of clusters was the predicted three for the di-/tri-cyclic as well as three for the C₃-/C₅-alkyl cannabinoid fractions (Figure 5A). The categories of the F_dicyclic values formed from the cluster analysis were congruent with those determined from the F_dicyclic: F_tricyclic frequency distributions (Figures 3B,C), with plants being categorized into low, intermediate and high F_dicyclic value classes (Figure 5B). For the F_C3 values, plants were also categorized into low, intermediate and high classes (Figure 5C), with the F_C3 clusters ranging between 0.43–22.81, 16.87–67.14, and 61.91–91.70%, respectively.

DISCUSSION

Plasticity of Alkyl Cannabinoid Composition

The quantity and quality of secondary plant metabolites are often attributed to a combination of genetic and environmental (G x E) factors (Bustos-Segura et al., 2017), with chemotypic plasticity associated with changing expression patterns in response to biotic and abiotic cues (Wink, 2003). Under environmentally uniform conditions we found that the di-/tri-cyclic as well as the C₃-/C₅-alkyl cannabinoid fractions were relatively stable throughout development, which is consistent with previous reports of C₃-alkyl cannabinoid composition from clonal (De Backer et al., 2012; Aizpurua-Olaizola et al., 2016) and seed propagated plants (Pacifico et al., 2008) grown in controlled environments. This suggests that the between-plant variation in cannabinoid quality observed within the diversity collection has a strong genetic influence independent of intragenerational environmental stimuli, and that the di-/tri-cyclic as well as the C₃-/C₅-alkyl cannabinoid chemotypes may have developed over longer periods via anthropogenic selective pressures and/or clonal adaptation. Indeed, intraspecific comparisons of Artemisia californica grown in a common environment together with precipitation manipulation treatments have shown limited plasticity in terpenoid quality, with compositional dissimilarity associated with source latitudinal distance (Pratt et al., 2014).

The between-plant alkyl cannabinoid chemotypic variation could have also been generated by the response of ecotypically distinct genotypes to a homogeneous environment.
cyclisation of the major di-/tri-cyclic cannabinoids have also been poorly correlated to THCA (Cascini et al., 2013) and CBDA proportions (Onofri et al., 2015), while the presence or absence of functional THCAS and CBDAS genes has been found predictive of cannabinoid quality (Weiblen et al., 2015). Given that THCA:CBDA cannabinoid proportions typically follow Mendelian inheritance (De Meijer et al., 2003), and that crosses between high C3-alkyl cannabinoid inbreeds and a high C5-alkyl cannabinoid clone form F1 progeny with distinct C3-/C5-alkyl cannabinoid chemotypes intermediate to the parents (De Meijer and Hammond, 2016), a predominant genetic basis for cannabinoid quality is unambiguous.

Recent discoveries in the genomic organization of secondary plant metabolism genes and associated transcriptional regulatory mechanisms may provide explanation for the stability of the di-/tri-cyclic as well as the C3-/C5-alkyl cannabinoid fractions. The occurrence of non-homologous secondary metabolite gene clusters has been well documented in a number of diverse plant taxa (Boycheva et al., 2014). Chromatin immunoprecipitation analysis in Arabidopsis thaliana has shown that the histone variant H2A.Z facilitates localized nucleosome opening and expression of contiguous thalianol as well as morphenal gene clusters, with independently formed clusters encoding product-specific oxidosqualene cyclases, cytochrome P450 enzymes and acyltransferases required for the synthesis of these triterpenoids (Nützmann and Osbourn, 2015). Despite limited characterization at all levels of gene cluster regulation, including analysis of promoter and cis-regulatory elements (Nützmann et al., 2016), evidence for the coordinated expressing of 43 secondary metabolic clusters has also been identified using the ATTED-II coexpression database (Aoki et al., 2016) in A. thaliana, Sorghum bicolor, Oryza sativa, and Solanum lycopersicum (Schläpfer et al., 2017).

It may be possible that the coordinated transcriptional regulation of non-homologous cannabinoid gene clusters limits expressional selectivity of cannabinoid pathway genes. This may result in increased stability of cannabinoid compositional homogeneity throughout development and limit variation in cannabinoid composition to heritable recombination events. While no direct observation of non-homologous gene clusters has yet been identified in Cannabis, evidence for tandem duplication of THCAS (McKernan et al., 2015) and potentially CBDAS (Onofri et al., 2015; Weiblen et al., 2015) as well as single gene transposition from long interspersed element-like (LINE-like) retrotransposons (Sakamoto et al., 2000) suggest that genomic reorganization mechanisms associated with metabolic gene cluster formation (Schläpfer et al., 2017) may have occurred. Completion of a fully annotated and chromosome-anchored genome assembly for Cannabis (Van Bakel et al., 2011; Vergara et al., 2016) may provide opportunities to elucidate the functional genomic architecture responsible for cannabinoid compositional stability. Functional characterization of alkyl-cannabinoid-determining loci may allow application of gene editing technologies, such as clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated9 (Cas9) (Alagöz et al., 2016), for development of elite chemotypes capable of producing alkyl cannabinoids beyond that of C3 or C5 configurations (Vree et al., 1972; Smith, 1997). Genetic enhancement and precise metabolic engineering of the alkyl pharmacophoric element could not only lead to therapeutic cannabinoid portfolio expansion (De Meijer and Hammond, 2016), but may also facilitate quality improvement of plant-based cannabinoid production systems (Potter, 2014; Chandra et al., 2017).

**Chemotypic Heterozygosity**

Heterozygosity at multiple chemotype-determining loci may account for a reduction of variance explained in the F\textsubscript{dicyclic}/F\textsubscript{tricyclic} values between vegetative and maturation stages in a subset of East Asian individuals. Allelism tests on progenies segregating for THCA and CBDA support a co-dominant B locus model, whereby the alleles encoding THCA and CBDA synthase govern THCA:CBDA cannabinoid proportions (De Meijer et al., 2003). DNA marker analysis of Cannabis chemotypes has shown that F\textsubscript{dicyclic}/F\textsubscript{tricyclic} values of ≥90% are associated with THCAS or CBDAS homozygosity, while intermediate chemotypes with F\textsubscript{dicyclic}/F\textsubscript{tricyclic} values of <90% are associated with THCAS and CBDAS heterozygosity (Welling et al., 2016a). In the THCAS:CBDAS heterozygote state, functional synthases are believed to compete for the substrates CBGA and CBGVA (Shoyama et al., 1984). The catalytic efficiency of THCA and CBDA synthases are reported to be dependent on alkyl side chain length (Shoyama et al., 1984), which suggests that metabolic fluxes of CBGA or CBGVA substrate within a THCAS:CBDAS heterozygote individual could lead to transitional changes in the F\textsubscript{dicyclic}/F\textsubscript{tricyclic} ratio.

To test whether the activity of THCA and CBDA synthase could be affected by CBGA or CBGVA substrates, we compared the F\textsubscript{dicyclic}/F\textsubscript{tricyclic} values within the C3-/C5-alkyl cannabinoid fractions in mature THCAS:CBDAS heterozygote plants (n = 20). Despite a wide range of F\textsubscript{dicyclic}/F\textsubscript{tricyclic} dissimilarity between the C3-/C5-alkyl cannabinoid fractions among genotypes, the F\textsubscript{c5} F\textsubscript{dicyclic}/F\textsubscript{tricyclic}: F\textsubscript{C3} F\textsubscript{dicyclic}/F\textsubscript{tricyclic} ratio was 1.44 (±0.34%). Interestingly, the individuals M01, M02, and M04 which in the truncated chemotypic distribution contributed to developmental F\textsubscript{dicyclic} variation, exhibited both the F\textsubscript{dicyclic} and F\textsubscript{C3} values close to parity at maturation, with M04 exhibiting F\textsubscript{dicyclic} and F\textsubscript{C3} values of 51.67% (±0.18%) and 54.41% (±0.22%), respectively. Given that these individuals are likely THCAS: CBDAS heterozygotes which can produce both C3- and C5-alkyl cannabinoid precursors, substrate flux above either THCA or CBDA synthases’ K\textsubscript{m} could result in substrate competition that affects the steady state concentration and time-dependent behavior of cannabinoid end products (Schäuble et al., 2013), resulting in the non-conformity of the di-/tri-cyclic cannabinoid fractions observed between vegetative and maturative stages.

**Genetic Regulation of Alkyl Cannabinoid Composition**

Despite the therapeutic importance of the cannabinoid alkyl side chain, the biosynthetic and genetic relationships responsible for alkyl homolog specificity remain poorly characterized in Cannabis. In the case of C3-alkyl cannabinoids, the prenylated
resorcinyl core and alkyl side chain are formed from the fatty acid starter unit hexanoic acid. This undergoes cytosolic acyl-activation (Stout et al., 2012) as well as polyketide formation by a tetraketide synthase (TKS) and olivetolic acid cyclase (OAC) complex forming the alkylresorcinol olivetolic acid (Gagne et al., 2012), prior to aromatic prenylation by geranyl-pyrophosphate:olivetolate geranyltransferase (GOT) (Fellermeier and Zenk, 1998) forming CBGA.

A similar mechanism, involving butanoic acid as a starter unit and the alkylresorcinol divarinic acid, is predicted for the synthesis of CBGVA. This is based on the functional characterization of recombinant alkylresorcinol synthases in the Poaceae plant family, which utilize acyl-CoA variously to form alkylresorcinol side chain homologs (Cook et al., 2010), as well as TKS (Taura et al., 2009) and GOT (Page and Boubakir, 2011) accepting butanoyl-CoA and a variety of aromatic substrates, respectively. However, the origin and synthesis of hexanoic and butanoic acid are unknown (Marks et al., 2009; Stout et al., 2012), while understanding the contribution of intracellular compartmentation, including metabolon constructs, on the channeling, selection and utilization of cannabinoid precursors, is incomplete. Moreover, the enzymatic promiscuity or specificity of OAC (Gagne et al., 2012) and GOT (Page and Boubakir, 2011) has not been examined with the predicted C3-alkyl cannabinoid intermediates. Nonetheless, it appears plausible that changes in the alkyl side chain originate prior to and possibly at polyketide formation, implying that multiple loci contribute to C3-alkyl cannabinoid composition.

Allelism tests suggest that an oligogenic or polygenic multi-locus $A^1$-$A^2$-$\ldots$ $A^n$ governs the C3-/C5-alkyl cannabinoid ratios in plants, although discontinuities in the C3-/C5-alkyl cannabinoid distributions of the available progeny were inadequate to form categorizations based on cannabinoid quality (De Meijer and Hammond, 2016). From the cluster analysis of within-plant variation, we identified three discrete FC3/FC5 categories (Figure 5C). As for the di-/tri-cyclic cannabinoid fractions (Figure 5B), the presence of three categories could indicate a monogenic model for C3-alkyl cannabinoid chemotypes, whereby allelic variation governing alkylresorcinol fatty acid starter unit availability or incorporation facilitates changes in the FC3/FC5 ratio. In a C3-/C5-alkyl cannabinoid monogenic model, small chemotypic differences between genotypes coupled with large individual variation within genotypic classes, could explain phenotypic continuity (Griffiths et al., 1999). However, the apparent absence of extreme individuals with FC3 values $\geq$90% within the sample population suggests the potential for additional categories, which would support an oligogenic or polygenic mechanism. In any case, the FC3/FC5 clusters identified are consistent with categorizations which can be expected within genetic resources of Cannabis and therefore offer utility in the selection and breeding of C3-alkyl cannabinoid genotypes.

As licit large-scale multi-billion dollar industries based on Cannabis emerge in the United States (Butsic et al., 2017), small incremental changes in the relative proportions of cannabinoids could have significant commercial and therapeutic implications for botanical drug development and manufacture (Potter, 2014; Chandra et al., 2017). Through selective inbreeding and hybrid clone selection, GW Pharmaceuticals, plc have reportedly achieved double- and triple-cross inbred plant lines with C3-alkyl cannabinoid proportions up to 96% (De Meijer and Hammond, 2016). In the current analysis we demonstrated a wide range of the C3-/C5-alkyl cannabinoid proportions within a relatively small subset of individuals from a single generation, which highlights the value of Cannabis ex situ conservation and characterization (Welling et al., 2016a). Comprehensive sampling of Cannabis genetic resources, both within and between accessions (Soler et al., 2017; Figure 3A), may make it possible to identify and select for pharmaceutically valuable chemotypes capable of reaching FC3 values $\geq$96%. However, it is uncertain whether the C3-alkyl cannabinoid fraction could match or exceed the C5-alkyl cannabinoid fraction in chemotypically extreme individuals. This may be affected by the lower molecular weight of C3-alkyl cannabinoid homologs which leads to a disproportionately reduced representation when comparing fractions/proportions derived from weight per weight concentrations.

**CONCLUSION**

The major alkyl cannabinoids of Cannabis were characterized across three developmental stages within a chemotypic diversity panel. Under controlled conditions alkyl cannabinoid composition was found to be stable throughout development. This suggests a strong genotypic influence on alkyl cannabinoid compositional variation and the potential for genetic enhancement of the alkyl pharmacophoric element. Further chemical and genomic characterization of Cannabis genetic resources may provide greater insight into the genetic mechanisms responsible for alkyl cannabinoid composition and provide novel opportunities for the genetic metabolic engineering and pharmaceutical diversification of plant derived alkyl cannabinoid chemotypes.

**AUTHOR CONTRIBUTIONS**

MW designed and performed the experiments and prepared the manuscript. LL provided contributions to conception and design of the research project, as well as development of analytical procedures, and provided detailed review and revision of the manuscript. CR performed statistical analyses and review and revision of the manuscript. OA provided background information and performed review and revision of the manuscript. GK provided substantial contributions to conception and design of the research project and performed detailed review and revision of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2018.01510/full#supplementary-material

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