

Jury Member Report – Doctor of Philosophy thesis.

Name of Candidate: Rim Gubaev

PhD Program: Life Sciences

Title of Thesis: Genetic association mapping for agronomically important traits in rapeseed and sunflower

Supervisor: Professor Philipp Khaitovich

Name of the Reviewer: Prof Dr Yalcin KAYA

I confirm the absence of any conflict of interest (Alternatively, Reviewer can formulate a possible conflict)	Date: 03-09-2022
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The purpose of this report is to obtain an independent review from the members of PhD defense Jury before the thesis defense. The members of PhD defense Jury are asked to submit signed copy of the report at least 30 days prior the thesis defense. The Reviewers are asked to bring a copy of the completed report to the thesis defense and to discuss the contents of each report with each other before the thesis defense.

If the reviewers have any queries about the thesis which they wish to raise in advance, please contact the Chair of the Jury.

Reviewer's Report

Reviewers report should contain the following items:

- Brief evaluation of the thesis quality and overall structure of the dissertation.
- The relevance of the topic of dissertation work to its actual content
- The relevance of the methods used in the dissertation
- The scientific significance of the results obtained and their compliance with the international level and current state of the art
- The relevance of the obtained results to applications (if applicable)
- The quality of publications

The summary of issues to be addressed before/during the thesis defense

Oil crops are the most important industrial crops in the world. Blacksea region has over 60% of world sunflower production and rapeseed is also major crop in the area. Russia is the leading country for both crops in the world. Thesis focused on most important crops for the country and also the most significant traits for both crops Mr Gubaev investigated in these valuable traits for sunflower and rapeseed within the framework of the PhD thesis research to determine potential benefits for breeders and researchers to develop better cultivars.

The author conducted a tremendous amount of research, which allowed him to reach the goal. His performed work has a major applicative value, especially evidenced in practical recommendations for future studies in this subject. Methodology used by the author is combined harmoniously both traditional and up-to-date methods. The obtained data are reliable. He reviewed many literatures and then he draws logical conclusions of his research results with comparing many previous studies. He performed very well and high level scientific publications too.

I included some minor corrections in thesis file.

Furthermore, I have some suggestions:

- In the abstract and conclusion parts, the author should mention some practical results of the conducted research, because there are no concrete results and suggestions for researchers who will work on same subjects in the sunflower and rapeseed as well as breeders how to utilize these findings and adapt their breeding programs.
- The author should mention especially for molecular results how to transfer these findings to formulate and facilitate selection process for classical breeding program based on study results.
- In some part of the thesis, some results especially molecular studies are not clearly mentioned for important traits. For instance, these used markers are polymorphic or not, it could be used for all genotypes and these are genotype dependent or not.

I considered and evaluated his PhD thesis “Genetic association mapping for agronomically important traits in rapeseed and sunflower”, and then I am fully agreeing that PhD thesis meets the high requirements for such kind of works and he deserves scientific title of PhD in the life sciences.

Provisional Recommendation

I recommend that the candidate should defend the thesis by means of a formal thesis defense

I recommend that the candidate should defend the thesis by means of a formal thesis defense only after appropriate changes would be introduced in candidate's thesis according to the recommendations of the present report

The thesis is not acceptable and I recommend that the candidate be exempt from the formal thesis defense

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GENETIC ASSOCIATION MAPPING OF AGRONOMICALLY
IMPORTANT TRAITS IN RAPESEED AND SUNFLOWER

Doctoral Thesis

By

RIM GUBAEV

DOCTORAL PROGRAM IN LIFE SCIENCES

Supervisor

Professor Philipp Khaitovich

Moscow – 2022

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I hereby declare that the work presented in this thesis was carried out by myself at Skolkovo Institute of Science and Technology, Moscow, except where due acknowledgment is made and has not been submitted for any other degree.

Candidate (Rim Gubaev)

Supervisor (Prof. Philipp Khaitovich)

Abstract

Oilseeds crops are the main source of vegetable oil production. Constant population growth, new industrial applications, as well as climatic change, require the selection of the high yield varieties that are adapted to changing populational requirements and unfavorable environmental conditions. Additionally, current breeding programs should meet the high-quality standards of the industry. One of the established approaches facilitating the breeding is marker-assisted and genomic selection, relying on modern genotyping techniques. Both approaches reduce the time and financial costs needed for plant evaluation in the field through the selection and monitoring of the preferred genotypes by means of genetic tests. One of the major steps that underlies both approaches is a genetic characterization of the mapping population followed by the identification of the genetic markers associated with the trait of interest. In Russia, the top three oilseed crops are sunflower, rapeseed, and soybean. Despite their substantial economic and food safety value, the application of genetic-information-based breeding approaches to these crops in Russia is currently lagging. This thesis focuses on the investigation of genetic markers associated with agronomically important traits in rapeseed and sunflower with the intent of enhancing marker-assisted rapeseed and sunflower breeding in Russia.

Publications

Main publications:

Gubaev, R.; Boldyrev, S.; Martynova, E.; Chernova, A.; Kovalenko, T.; Peretyagina, T.; Goryunova, S.; Goryunov, D.; Mukhina, Z.; Ben, C.; Gentzbittel, L.; Khaitovich P.; Demurin, Y. (2022). Genetic mapping of loci involved in oil tocopherol composition control in Russian sunflower (*Helianthus annuus* L.) lines. G3: Genes, Genomes Genetics, 12, 4, jkac036. <https://doi.org/10.1093/g3journal/jkac036>

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List of abbreviations

AFLP – Amplified fragment length polymorphism

ANOVA – Analysis of Variance

BC – Back cross

bp – base pair

CIM – Composite interval mapping

cMLM – Compressed mixed linear model

CMS – Cytoplasmic male sterility

DH – Doubled haploid

FA – Fatty acid

GBS – Genotyping by sequencing

GC – Gas chromatography

GS – Genomic selection

GWAS – Genome-wide association study

IMI – Imidazolinone

kb – kilobase

LAMP – Loop-mediated isothermal amplification

LD – Linkage disequilibrium

LOD – Logarithm of the odds

MAGIC – Multi-parent advanced generation inter-cross

MAS – Marker-assisted selection

Mb – Megabase

MLM – Mixed linear model

MPPs – Multi-parental populations

MS – Mass spectrometry

NAM – Nested association mapping

NGS – Next generation sequencing

PC – Principal component

PCA – Principal component analysis

PCR – Polymerase chain reaction
PVE – Phenotypic variation explained
QTL – Quantitative trait loci
R gene – resistance gene
RAD – Restriction-site associated DNA
RAPD – Random amplified polymorphic DNA
RFLP – Restriction fragment length polymorphism
RILs – Recombinant inbred lines
SIM – simple interval mapping
SLAF – Specific-locus amplified fragment sequencing
SNP – Single nucleotide polymorphism
SSR – Simple sequence repeat
SU – Sulfonylurea
TLC – Thin-layer chromatography
VIR – N.I.Vavilov Research Institute of Plant Industry
VNIIMK – Pustovoit All-Russia Research Institute of Oil Crops

Chapter 1. Introduction

Starting from the neolithic period people started to domesticate wild plants turning them into crops, thus securing a controlled food supply. From then on plant breeding relying on hybridization and selection continued fundamentally unchanged till the introduction of genetics. The genetics concepts entered the plant breeding field with the rediscovery of Mendel's Laws of Heredity at the end of the 19th century. Within the 20th century, increasing understanding of genetic mechanisms led to important innovations in plant breeding. These innovations included spontaneous mutagenesis, relying on the application of radiation and chemicals, which facilitated the creation of plants with the traits lacking in the natural populations. In addition, directed mutagenesis based on the *Agrobacterium*-mediated transformation and gene gun technique were applied to transfer genes introducing novel agronomically important traits. The development of the molecular genetics approaches and DNA sequencing technologies led to another technological innovation called marker-assisted selection implementing genetic markers whose allelic states, in turn, may indicate the presence or absence of a specific trait. This technology allowed breeders to accelerate the breeding process by replacing lab or field phenotypic assessment with genetic testing and optimizing the breeding schemes, reducing time and financial expenditure. At the beginning of the 21st century, the other two technological breakthroughs emerged: genetic editing techniques allowing the introduction of site-specific mutations in the genes of interest and high throughput genotyping. The high throughput genotyping of crops facilitated studies of genetic diversity and improved prediction of complex polygenic traits providing a basis for improved genomic selection.

All above mentioned approaches and technologies are nowadays applied in the broad spectrum of crops including oilseeds – the major source of vegetable oil, an important part of our diet. In addition to their dietary role, vegetable oil and oilseed by-products are used to produce biofuels, chemicals and serve as fodder for animals. The top five mostly produced oilseeds worldwide include soybean, rapeseed,

sunflower, peanut, and cottonseed. Due to the constant population growth, the demand for vegetable oil permanently increases. Additionally, increasing food quality standards, food and chemical industry requirements, and ecological and climate alterations require constant improvement of specific oilseed traits, such as productivity, composition and quality of the oil, and resistance to pathogens and unfavorable environmental conditions. All of these challenges could be met more timely by the implementation of genetics-based technological approaches.

In Russia, the most produced oilseed crops are sunflower, rapeseed, and soybean. Importantly, Russia is one of the world's leaders in sunflower production. Despite that, modern genetic technologies are not broadly used by scientific organizations and breeding companies. As a consequence, two problems arise. First, Russian oilseed germplasm is poorly characterized in terms of genetic diversity. Second, the implementation of genetic modification approaches and marker- and genome-assisted breeding is not a common trend. This impedes the creation of new competitive varieties meeting market demands. As a consequence, Russian oilseed producers often use imported seed material due to its higher quality, rooted in more efficient breeding strategies involving genetics-based technological solutions.

To improve this situation, several attempts are now made with an aim of supporting the introduction of novel genetic and biotechnological innovations in Russian scientific institutions and breeding companies. The present work is part of the bigger initiative supported by the Ministry of Science and Higher Education aimed at developing marker- and genomic-assisted approaches in oilseed crop breeding for providing food security in Russia. Within this initiative, the Skolkovo Institute of Science and Technology (Skoltech) collaborated with leading organizations involved in germplasm preservation and breeding of oilseeds, namely, the Vavilov Institute of Plant Industry (VIR), Pustovoit All-Russia Research Institute of Oil Crops (VNIIMK), and Agroplasma breeding company. The present thesis focuses on the association mapping of agronomically valuable traits in rapeseed and

sunflower. Additionally, a basic analysis of genetic diversity in rapeseed and sunflower germplasms from the above-mentioned organizations was performed.

The aim of the present study was to map agronomically important traits in rapeseed and sunflower. The following objectives were set within this aim:

- 1) Find the genetic associations for glucosinolate content by analyzing the rapeseed diversity panel from the VNIIMK.
- 2) Find genetic associations with the tocopherol composition and oleic acid content of sunflower oil by analyzing experimental crosses generated by the VNIIMK.
- 3) Find genetic associations with seed and husk size, seed to husk ratio using the diversity panel from VNIIMK, VIR and Agroplasma.
- 4) Characterize the genetic diversity of the rapeseed collection from VNIIMK.
- 5) Characterize the genetic diversity of the Russian sunflower collection from VNIIMK, VIR, and Agroplasma.

Chapter 2. Literature review

2.1 Marker-assisted and genomic selection as tools for plant breeding

Plant breeding is a set of methods for creating plant varieties and hybrids with the desired properties with an aim of increasing the yield and quality of crops. Starting from the neolithic revolution, people began to domesticate plants, select and reproduce ones with the desired properties. For a long time plant breeding was performed empirically without any strong scientific background. Systematic plant breeding based on hybridization and selection began to be possible in the early 20th century after the rediscovery of Mendel's laws (Lee et al., 2015).

The idea of using markers, associated with certain traits that may help to facilitate the selection of prospective progeny appeared in the first half of the 20th century. The first markers were morphological traits, such as seed coat color, associated with the seed size in a common bean (Sax, 1923), or flower color associated with flowering time in pea (Rasmusson, 1935). In addition to morphological markers, cytological and biochemical ones were developed throughout the 20th century. To date, however, the majority of markers of traits are DNA-based (Nadeem et al., 2018).

The concept of marker-assisted selection (MAS) based on the DNA markers became popular in the 1980s, partially due to the development of laboratory techniques that allowed the identification of genetic polymorphisms among individuals in the population, some of which were related to the specific traits of interest (Ben-Ari & Lavi, 2012). The main idea was the following: not all genetic polymorphisms are associated with a trait of interest; however, some of them could be associated with monogenic or polygenic traits controlled by quantitative trait loci (QTL) and thus may substitute field testing with predictions based on the markers' states (Smith & Simpson, 1986; Soller & Beckmann, 1983). The marker-assisted selection is based on the concept of linkage disequilibrium (LD), a non-random

association between alleles at a certain region on a chromosome affected by different factors including genetic linkage, selection, mutation, non-random mating, population structure, genetic drift and recombination. Thus a genetic marker of a trait could represent a causal mutation in a gene that affects the phenotype (rare case) or it could be linked to the allele of the gene affecting the phenotype by LD (Mackay & Powell, 2007). MAS is commonly used to mark traits associated with the resistance to pathogens and R-gene pyramiding, resistance to herbicides, ability to restore fertility etc., It should be noted that most of these traits are mono-/oligogenic (Bulos, 2013; Melchinger, 1990; Ramalingam et al., 2020).

The modern marker-assisted selection is based on the implementation of genetic markers that could be of a different kind. To date, five more common groups of genetic markers could be highlighted: restriction fragment length polymorphism (RFLP), random amplified polymorphic DNA (RAPD), amplified fragment length polymorphism (AFLP), Simple sequence repeats (SSR) and single nucleotide polymorphisms (SNPs). The first three marker systems (fragment-based polymorphisms) were the first ones to be applied widely in the 1980s and 1990s: RFLP and AFLP markers are based on the point mutations affecting the fragment restriction, while RAPD is based on point mutations that affect primer annealing (Ben-Ari & Lavi, 2012). All these markers do not require prior information about sequence but require gel electrophoresis for the detection, additionally, these markers are not so spread within a genome and to date are rarely used. Quite popular SSR markers represent short (2-6 bp long) repeated sequences that are widely distributed among plant genomes. The main advantage of these markers is a high level of polymorphism i.e. one marker can distinguish up to more than ten alleles (Kalia et al., 2011). The most commonly used markers nowadays are SNPs. The main advantage of the SNP markers is that these kinds of markers occur at higher rates through the genomes compared to the abovementioned markers which in turn makes it possible to find more genetic associations for traits of interest, especially polygenic ones. SNPs markers became widespread at the beginning of the 21st century partially due to the

rapid development of the high-throughput genotyping approaches, namely DNA microarrays and next-generation sequencing whose implementation towards the genetic diversity assessment is mostly concentrated on the detection of the SNP markers (Rasheed et al., 2017).

The ability to identify a high amount of genetic markers led to the development of the novel branch of marker-assisted selection called genomic selection (GS) at the beginning of the 21st century (Meuwissen et al., 2001). Unlike MAS which mostly concentrates on the one and/or several QTL associated with mono-/oligo genic traits with high heritability, GS relies on the hundreds or thousands of genetic markers, (usually SNPs or SSR) with small effects (Crossa et al., 2017). This allows the estimation of genome-estimated breeding values (GEBVs) of a certain individual and to select candidates for the breeding program (Nadeem et al., 2018). To date, several attempts have been made to facilitate the development of GS-based breeding programs, most of them were done on crops (wheat, maize, rice), and the key traits to improve included yield and resistance to unfavorable environmental conditions (Bhat et al., 2016).

The advantage of the MAS and GS results from the fact that the vast majority of the traits of interest are not easily assessed, and usually need additional laboratory testing (resistance to pathogens, metabolite content assessment, etc.) or field trials (yield, resistance to unfavorable environmental conditions, etc.). Thus the implementation of MAS and GS allows for reducing the time and financial costs of breeding (Ben-Ari & Lavi, 2012). It also should be noted that in recent years startups like NRGene, NeoGene, GeneCove e.t.c., are emerging and suggest a service in MAS and GS for breeding and seed companies with the aim of implementing genetic markers to facilitate new varieties' development.

Summarizing the information reviewed above it could be concluded that marker-assisted and genomic selection is now widely applied with the aim of obtaining new varieties at a higher rate with lower costs. It should be noted that the development of marker-assisted breeding approaches is based on the scanning for the

genetic markers of the agronomically important traits with subsequent development of genetic tests as well as marker panels.

2.2 Approaches for finding genetic markers and developing genotyping solutions

A basic procedure that underlies marker-assisted selection is scanning for the genetic markers for the traits of interest. Search for genetic markers consists of the following steps: 1) selection of the association panel, 2) phenotyping, 3) genotyping, 4) association mapping and 5) marker validation.

The scanning for the genetic markers usually starts with the creation and analysis of the population. In general, the two most common types of mapping populations are currently used to perform genetic mapping. The first one includes the diversity panels, which are usually formed by sampling the gene banks' accessions of phenotypically and genetically contrasting lines that may include wild, domesticated, landraces and modern ones (Brachi et al., 2011). The second type involves artificial populations obtained by means of experimental crosses. Artificial populations, in turn, may be classified in bi-parental including F₂, double haploid (DH) backcross (BC), and recombinant inbred lines (RILs) populations and multi-parental populations (MPPs), including nested association-mapping populations (NAMs), multi-parent advanced generation inter-cross (MAGIC). Bi-parental populations are based on the crossing of two parents contrasting in the trait of interest followed by a subsequent rounds of selfing (F₂, RILs), backcrossing (BC), or production of doubled haploids (DHs). Multi-parental populations include several parents: NAM consists of a series of biparental populations having one common founder as a parent. MAGIC has a simple funnel breeding design where several unrelated parental pairs (4-16) are crossed (Scott et al., 2020).

The advantage of the diversity panels is the higher resolution of association mapping. The analysis of these populations is based on the detection of associated regions generated by historical recombination events. Thus it is possible to detect loci

associated with traits in the narrow genomic regions (Mohammadi et al., 2020). The advantage of the bi-parental populations is the ability to study traits with low natural diversity, or traits artificially introduced by spontaneous mutagenesis (Oladosu et al., 2016). Additionally, only two generations are required to produce a mapping population. However, the mapping resolution remains suboptimal due to a low number of recombination events (Arrones et al., 2020). Multi-parental populations allow for increasing the mapping resolution compared to bi-parental populations. Further, it is possible to artificially increase the frequency of the rare alleles with an aim of detecting rare variants associated with the traits of interest. The disadvantage of MPPs is the substantial time needed to produce crosses and selfing to create a mapping population (Pascual et al., 2015).

The next step preceding the marker scanning is phenotyping, which is to some extent more important than the subsequent genotyping since the accuracy and replication of the phenotype data affect the prediction accuracy of the identified markers. Additionally, phenotyping costs are larger than the genotyping ones as genotypic data for a mapping population is obtained once and can be used to map several traits of interest. Thus, the collection of phenotype data might create a bottleneck in association studies due to the time and labor costs needed for its collection (Reynolds et al., 2019). Another important aspect of phenotyping is the collection of the phenotype under different climatic/year conditions and the usage of standardized protocols. Trials in several geographically distant fields and/or trials across different years facilitate the identification of a trait variance explained by the genotype x environment (GxE) and improve the accuracy of the prediction models for GS and markers identified by GWAS or QTL mapping (El-Soda et al., 2014). Standardized protocols, on the one hand, facilitate the collection and reusing of the phenotype data and on the other hand improve the exchange of the data among the plant breeding community (Pieruschka & Schurr, 2019).

An important procedure related to phenotyping is the collection of relevant environmental data, which mostly includes temperature, solar radiation, and

precipitation over a specific vegetation period. Recent trends in this direction lead to the development of so-called environmentomics which in addition to climatic data assumes the soil composition and structure. Microbiome data was also suggested to be included (Oyserman et al., 2021). All the above-mentioned information could be used to improve the accuracy of the yield and phenotype prediction models, especially for crops grown in contrast environmental conditions (Rogers et al., 2021).

The final step before the genetic association analysis is genotyping: detecting genetic polymorphisms in studied populations. The genotyping of the plants for scanning for the markers started in the 80s with low-throughput RFLP markers systems (Tanksley et al., 1989) and nowadays is mostly based on more occurring throughout the genome marker, namely SSRs and SNPs. Starting from the 2000s, following the deciphering of the plant genomes, DNA microarrays started to be used widely in order to genotype plant species including major crops like maize, rice and soybean (Rasheed et al., 2017).

The advantage of the DNA microarrays is related to their standard design and their ability to capture a significant amount of genetic diversity existing in specie compared to fragment-based polymorphisms analysis since the arrays are done within the consortium projects that usually analyze several germplasm collections of different origins (Morales et al., 2020; Unterseer et al., 2014). The power of the DNA-microarrays is restricted to the number of species for which such technology is available and the relatively high cost of the array's design and production for a small number of samples (Christiansen et al., 2021). In addition to DNA microarrays, high-throughput sequencing technologies are currently gaining popularity for plant genotyping due to their rapid development and constant reducing DNA sequencing cost (Egan et al., 2012). Although whole-genome sequencing data gives the opportunity to obtain complete information about causative and linked polymorphisms, it remains expensive to be obtained and computationally heavy to analyze. This, lead to the development of reduced genotyping approaches like restriction-site associated DNA sequencing (RAD-seq), specific-locus amplified

fragment sequencing (SLAF-seq), genotyping-by-sequencing (GBS) that are based on the sequencing of the restricted DNA-fragments (Baird et al., 2008; Elshire et al., 2011; X. Sun et al., 2013). These techniques allow for reducing the plant genome complexity overwhelmed with repetitive elements, as well as reducing genotyping costs and time for analysis.

Different types of examined population designs require different types of analysis to identify genetic markers. For bi-parental populations (BC, F2, RILs, DHs), the construction of the genetic maps based on the recombination fractions (RF) and LOD scores calculated between the pairs of the markers is required in order to reconstruct the true order of each marker in each linkage group (C. Zheng et al., 2019). Next, several approaches could be applied to scan for the associations, including single marker analysis which considers each marker as an independent observation, simple interval mapping (SIM), and more recently commonly used composite interval mapping (CIM) scans markers in the genetic intervals of linked markers and controls the effects of the QTL on other intervals or chromosomes on the tested QTL (Broman et al., 2003). For diversity panels, the genome-wide association studies approach (GWAS) is used (Brachi et al., 2011). Initially, GWAS was based on the single marker ANOVA that tests whether the marker is associated with the trait of interest (Bush & Moore, 2012). Such a naive approach does not take into account relatedness among individuals which may bias the results, since polymorphisms present at a higher frequency in a certain population could be falsely associated with the phenotype of interest of this population happened to be present at a higher frequency in a group expressing this phenotype (Tibbs Cortes et al., 2021). More recent GWAS methods are based on the linear models among which the mixed linear model (MLM) is widely used since it allows to account for the relatedness at two levels: kinship (K) and population structure (Q) (Yu et al., 2006). Kinship matrix (K) is used to control the random effects of genetic background, while the Q matrix explaining sample stratification, usually assessed by the principal component analysis or genetic clustering algorithms implemented in software like ADMIXTURE or

STRUCTURE, helps to control fixed effects of the population structure (Tibbs Cortes et al., 2021). In addition to the MLM, more than twenty additional approaches that are also based on the mixed models were developed in recent years mainly with the aim of reducing the computational demand and increasing the statistical power. Among them, SUPER (Settlement of MLM Under Progressively Exclusive Relationship), FaST-LMM (Factored Spectrally Transformed Linear Mixed Models) and FarmCPU (Fixed and random model Circulating Probability Unification) remain popular for trait mapping in plants (Gupta et al., 2019). Besides the family of linear model-based methods that often assume on and/or few markers at the time, Bayesian methods that use prior information on marker effects and phenotypes to estimate all marker effects simultaneously started to gain popularity. The main advantage of these methods is that the power of detecting associations is not inversely correlated to the number of polymorphisms as in the case of the application of multiple testing correction applied for linear model-based methods (Fernando & Garrick, 2013).

Regarding the MPPs several approaches could be highlighted. On the one hand, it is possible to apply linear models, as in the case of GWAS, for the diversity panels (Gangurde et al., 2020; Islam et al., 2016; Nice et al., 2016). On the other hand, since the parental genotypes are by default accessible for the NAM and MAGIC populations it is possible to construct a consensus genetic map for the studied population based on the information on recombination events (€ Zheng et al., 2019) and apply interval mapping approaches used for bi-parental populations that will increase the accuracy of the mapping (Bu et al., 2021).

After the candidate genetic markers ~~are found~~, they undergo validation based on the plant material used for breeding (Xu & Crouch, 2008). For this purpose, local genotyping tests, such as allele-specific PCR assays followed by the PCR product detection in an agarose gel, are commonly used. Then the predicted effects of these putative markers are compared to the test sample phenotype (L. Li et al., 2013; W.-C. Zhou et al., 2003). Additionally, such validation could be performed across different

environments and populations in order to assess the stability of the genetic markers (Knoll & Ejeta, 2008).

The next step after the genetic markers associated with the traits of interest have been verified based on the plant material used for the breeding and/or independent plant samples, is the development of a genotyping solution for routine material assessment. Here several technologies could be applied for the assessment of mono- and oligo-/poly-genic traits. Allele detection assays based on allele-specific PCR, for example SNP type (Fluidigm), KASPar (KBiosciences) and TaqMan (Life Technologies) could be used to predict the presence or absence of mono-/oligo-genic traits (Broccanello et al., 2018; Seeb et al., 2011). Notably, all the above-mentioned technologies require lab equipment. This obstacle is circumvented by the rapid development of isothermal PCR, like LAMP (loop-mediated isothermal amplification), which enables rapid phenotype prediction in the field (Prasannakumar et al., 2021; D. Zhou et al., 2016). For oligo-/poly-genic traits, alternative solutions like MALDI-based genotyping and low-density SNP-arrays are commonly applied (Pattemore et al., 2010; X.-L. Wu et al., 2016). The rapid development of the NGS further led to the emergence of the low-density NGS-based panels, for example, GT-seq (Genotyping-in-Thousands) technology, allowing multiplexing of sequence-specific regions carrying polymorphisms of interest (Campbell et al., 2015).

It could be concluded that scanning for the genetic markers requires a complex methodological pipeline including the choice of a suitable population design for the trait mapping, followed by a selection of optimal phenotyping and genotyping approaches. The collected data is then processed with an aim of scanning for candidate markers. Markers are then subjected to testing in different environmental and/or genetic backgrounds. All these steps lead to the creation of the marker panels, which are then used to reduce the financial and time costs for breeders in the context of marker-assisted breeding.

2.3 Sunflower and rapeseed: past and recent breeding trends

Rapeseed and sunflower are among the top three oilseed crops grown worldwide (Attia et al., 2021; Vinnichuk et al., 2019). These crops are grown mainly with the purpose of vegetable oil production. Besides vegetable oil, rapeseed and sunflower are used to produce raw materials for biofuels and the chemical industry (Embaye et al., 2018). Additionally, byproducts of oil production, namely, rapeseed oil cakes and sunflower meals are used to feed the animals (de Oliveira Filho & Egea, 2021; Walker & Booth, 2001).

Rapeseed

Rapeseed (*Brassica napus*) as a plant species appeared due to a spontaneous hybridization between two other *Brassica* species – cabbage (*B. oleraceae*) and turnip (*B. rapa*). According to phylogenetic analysis this event happened from 7500 to 12,500 years ago (Chalhoub et al., 2014) leading to allopolyploid *B. napus*. Domestication of rapeseed began in South and Southeast Asia around 6000 years ago (Snowdon et al., 2007). In Europe rapeseed was documented to be cultivated around 400 years ago (Gomez-Campo, 1999). Importantly, there are three ecotypes of rapeseeds: winter and semi-winter (lacking the frost hardiness) that are sown in the late autumn, and spring ecotype which is shown in early spring. Such plasticity has contributed to the widespread rapeseed cultivation across the world from Canada to Australia (D. Wei et al., 2017).

Significant changes in rapeseed breeding appeared throughout the second half of the 20th century. In the 1970s the first source of cytoplasmic male sterility (CMS) was discovered, making hybrid breeding for this species possible. However, the stable CMS source was transferred later from radish (*Raphanus sativus*) to rapeseed (Rousselle & Bregeon, 1982). Heterosis was reported for rapeseed only in the late 1980s (Brandle & McVetty, 1990). Next, substantial progress was made by Canadian breeders in the late 1970s – their efforts led to the development of new cultivars with low erucic acid and glucosinolates content in the oil (Mag, 1983). Lately, these traits were introduced in European germplasm (Przybylski et al., 2005). These

modifications are important because high erucic content leads to myocardial lipidosis – increased triacylglycerol accumulation affecting the contractility of the heart (J. Alexander et al., 2016). In turn, glucosinolates negatively affect the liver and kidney in animals (Walker & Booth, 2001) and contribute to bitter taste across the *Brassicaceae* family (Wieczorek et al., 2018). In addition to these traits, many efforts were made toward the selection of the high-oleic varieties of rapeseed in the 1990s, since oleic acid is considered optimal both for cooking and biofuel production (Spasibionek et al., 2020).

A more recent direction in rapeseed breeding is disease resistance. Three diseases mostly damaging rapeseed include stem rot (*Sclerotinia sclerotiorum*), blackleg disease (*Leptosphaeria maculans*) and clubroot disease (*Plasmodiopora brassicae*). For clubroot and blackleg there are qualitative disease resistance genes (R genes), thus the breeding for resistance mostly consists of scanning of the R gene in diversity panels and introduction to germplasm (Ton et al., 2020). For stem rot the disease resistance is quantitative and thus the breeding for resistance is more challenging (Z. Wang et al., 2019). Other important directions in rapeseed breeding related to the harvest preservation and effectiveness of harvesting include improvement of lodging resistance and pod shattering. Lodging resistance affects the uneven distribution of the light and the effectiveness of mechanical harvesting (C. N. Miller et al., 2018) while pod shattering is associated with yield loss during vegetation and harvesting as well (Kuai et al., 2016). As climate change significantly affects the sowing area for rapeseed (Jaime et al., 2018), several attempts and suggestions were proposed to diminish negative climatic effects. One of the breeding targets is the reduction of the flowering time as the rapeseed is sensitive to drought while transiting from the flowering stage to silique development (M. Zhu et al., 2016). Additionally, breeding for drought tolerance is also needed to reduce yield loss under non-optimal conditions (Khanzada et al., 2020).

Sunflower

Originally sunflower was domesticated by native North Americans around 4000 years ago (Crites, 1993). As an oilseed crop it was cultivated from the beginning of the nineteenth century in Russia. Significant progress in sunflower breeding was made by Soviet academician V. S. Pustovoit who developed open-pollinated varieties with increased oil content in the late 1950s (Terzić et al., 2020). After the discovery of a source for cytoplasmic male sterility (CMS) and the corresponding fertility restorer gene (Leclercq, 1971), sunflower hybrid breeding became widespread starting in the late 1970s (Dimitrijevic & Horn, 2018). Further, in the late 1990s, genes for imidazolinone and sulfonylurea herbicide tolerance were identified in the wild populations of sunflowers (Al-Khatib et al., 1998).

To date several trends in sunflower breeding could be highlighted. The first one is resistance to pathogens, including those that cause sunflower rust, downy mildew, sclerotinia white mold, etc. (Jocic' et al., 2015; Rauf et al., 2020; Škorić, 2016). The high interest for Eastern Europe and Russian breeders is represented by breeding for resistance to the broomrape (*Orobanche cumana*) – a parasitic plant that evolves rapidly and significantly affects yield (Lukomets et al., 2021; Škorić, 2016). Another important direction of sunflower breeding that is specific to Eastern Europe is breeding for large-seeded varieties (Hladni & Miladinović, 2019; Lukomets et al., 2021). Such varieties are called confectionary and are used to make edible seeds for direct consumption or to produce snacks (Hladni & Miladinović, 2019). The third large direction of breeding focuses on the oil content, composition and quality. The total oil content was increased by V.S. Pustovoit from ~25% to ~50% of seed mass. Now, modern breeding programs, especially the ones relying on hybrid breeding, are aimed to maintain and improve this complex trait using genomic prediction models (Mangin et al., 2017). For frying oils the priority is given to the high oleic acid oils, since this monounsaturated fatty acid increases oil resistance to thermal oxidation. Another direction of oil content selection aims to create high stearic acid sunflower for margarine and vegetable ghee manufacturing (Serrano-Vega et al., 2005).

Additionally, the tocopherol composition of the seeds is of high interest for oil quality improvement, since tocopherols demonstrate vitamin E activity and increase the resistance of oil to oxidation, especially in combination with high oleic acid content (Rauf et al., 2017).

A novel trend in sunflower breeding is drought resistance, as current climate changes may significantly affect future sunflower productivity. Although sunflower is a low-to-medium drought sensitivity crop, increased temperatures reduce the total oil content, total yield, and protein level (Hussain et al., 2008; Rondanini et al., 2003). Thus, breeding efforts are currently made, including the consortium level such as SUNRISE (~~SUNflower~~ Resources to Improve yield Stability in a changing Environment), with the main goal of making varieties that demonstrate high yield reproducibility across different environmental conditions (Debaeke et al., 2017).

Thus the breeding of both crops was aimed toward the improvement of the oil composition and yield quality for food and industry needs. As recent climate changes started to affect the yield, new breeding targets include resistance to unfavorable environmental conditions. Additionally, climatic changes accelerate pathogen evolution creating the need to select variants resistant to emerging pathogens.

2.4 Genetics of agronomically important sunflower traits

Being an industrially important species, agricultural and wild sunflower varieties are well characterized in terms of genetic diversity. The first genetic map of the sunflower was developed in the early '90s (Berry et al., 1995; Gentzbittel et al., 1995; Rieseberg et al., 1993) and its genome was sequenced using several approaches (Badouin et al., 2017). Accordingly, genotyping DNA microarrays were developed for this species. All of the above-mentioned features make a sunflower an appropriate object for marker scanning and subsequent application of marker-assisted selection approaches (Jocković et al., 2021). As a result, many important traits have already been dissected in terms of genetics for sunflowers.

To date, researchers uncovered more than 20 sources of cytoplasmic male sterility in *Helianthus* genus by experimental crosses (Terzić et al., 2020). However, only for two cytoplasms CMS-PET1 and CMS-PET2 derived from *H. petiolaris* the respective restorer genes have been identified (Z. Liu et al., 2012; Sajer et al., 2020, p. 2). For CMS-PET1 several germplasms were successfully scanned with an aim of finding the restorer genes (*Rf*) for CMS-PET1. It was demonstrated that loci associated with *Rf* genes mapped to chromosomes 2, 6, 12, 13, and 7 (Ma et al., 2021). Recently, *Rf-PET2* restoring CMS-PET2 was mapped on chromosome 13 (Sajer et al., 2020). For sunflower, as for other crops, these loci were demonstrated to carry genes encoding proteins carrying pentatricopeptide repeat (PPR) motifs (Goryunov et al., 2019; Horn et al., 2019; Ma et al., 2021; Sajer et al., 2020).

Weeds affect sunflower yield, thus the application of herbicides is economically important. For sunflower, two types of herbicides, namely sulfonylureas (SU) and imidazolinones (IMI) are applied. Both target acetohydroxyacid synthase key enzymes for branched amino acid biosynthesis encoded by *Ahsl* genes (Kaya et al., 2012). Herbicide-resistant plants were found among wild populations of the sunflower (Al-Khatib et al., 1998; White et al., 2002). Additionally, IMI-resistant plants were obtained based on spontaneous mutagenesis (Sala et al., 2008). Using mapping populations three herbicide-resistance loci and their corresponding genetic markers (SSRs, CAPS and SNPs) were identified and validated: *AhasII-1* (chromosome 9) determining moderate tolerance against both herbicides; *AhasII-2* (chromosome 6) conferring tolerance to SU; and *AhasII-3* (chromosome 2) determining resistance to IMI (Bulos, 2013; Kolkman et al., 2004).

There are more than 20 pathogens harming sunflower crops. Among them, the top three diseases leading to significant yield losses are caused by *Plasmopara halstedii* (downy mildew), *Orobancha cumana* (sunflower broomrape), *Sclerotinia sclerotiorum* causing white mold (Rauf et al., 2020; Virányi, 2008). More than 30 resistance genes to *P. halstedii* damaging to sunflower leaves have been identified in sunflower and in its wild relatives (Qi et al., 2019). However, only several of them

were mapped genetically to loci on chromosomes 1, 2, 4, 8, 13 and the corresponding genetic markers were identified (Dimitrijevic & Horn, 2018). Broomrape is a parasitic plant that feeds on the sunflower's root system leading to the reduction of leaf and flower head size. Broomrape is the most harmful sunflower pathogen in Eastern Europe (Škorić et al., 2021). To date, eight races (A-H) of *O. cumana* have been identified. Several genes of vertical resistance to broomrape have been identified and mapped as well. The most recently identified resistance gene (and corresponding SNP marker) was localized to chromosome 4 for race G by analyzing an experimental cross of resistant and susceptible sunflower lines (Martín-Sanz et al., 2020). Similarly, several studies reported lines carrying resistance to the race H. However, the genetic markers associated with this trait were not identified. Additionally, new potential broomrape races that overcome resistance to race H are emerging (Škorić et al., 2021). It should be noted that in addition to vertical resistance genes several QTLs that control sunflower resistance at different developmental stage of the broomrape were also identified (Louarn et al., 2016). Unlike the *P. halstedii* and *O. cumana*, resistance to *S. sclerotiorum*, a pathogen damaging all parts of the plants including the flower head, demonstrates mostly polygenic inheritance with narrow-sense heritability of up to 60% (Zubrzycki et al., 2017). Using F2 mapping populations over 15 potential QTLs associated with resistance to *S. sclerotiorum* were identified in sunflower (Micic et al., 2004). Of them, three SSR markers were validated using independent populations (Micic et al., 2005). In further studies, the utilization of the high-density SNP arrays could improve the accuracy of the *S. sclerotiorum* resistance predictions leading to a more rapid emergence of pathogen-resistant variants by genomic selection (Livaja et al., 2016; Talukder et al., 2021).

Sunflower total oil content ranges from 40 to 50% of dry seed mass in industrial varieties. Further, it was demonstrated that despite the complex genetic control of this trait, the estimated heritability was more than 80 percent (Bachlava et al., 2010). Several studies identified more than five QTLs controlling total oil content

explaining up to 51% of the trait variance (Bachlava et al., 2010; Merah et al., 2012). Application of the large-scale genotyping allowed to apply genomic selection approaches and to develop models for oil content prediction in hybrids, demonstrating high prediction accuracy of up to 78% (Mangin et al., 2017).

Genetic control of oleic acid content is one of the best-studied features, mainly due to the importance of this fatty acid to the food industry. Initially, it was assumed that the high oleic acid content is fully controlled by the dominant *Ol* gene located on chromosome 14 that encodes fatty acid desaturase FAD2 – an enzyme converting oleic acid to linoleic acid and explaining up to 86% of phenotypic variance (Lacombe & Bervillé, 2001). The first genetic markers associated with *Ol* were RAPD and RFLP were found by analyzing experimental crosses (BC and F2) (Dehmer & Friedt, 1998; Lacombe & Bervillé, 2001). Additional studies performed on the contrast F2 populations revealed additional minor effect loci associated with oleic acid content located on chromosomes 6, 8, and 9 and explaining up to 12% of the phenotypic variance (Premnath et al., 2016; F. Zhou et al., 2018). Besides oleic acid, several studies have been performed with an aim of mapping loci associated with other important fatty acids. For palmitic acid and stearic acids, which are of high importance for lubrication production and deep-fat frying, genetic markers were identified by experimental crosses analysis. QTL analysis revealed RFLP markers associated with the major effect locus controlling stearic acid content on chromosome 1 carrying a gene encoding stearyl-ACP desaturase and explaining up to 80% of phenotypic variance (Pérez-Vich et al., 2002). For palmitic acid locus affecting its content was mapped on chromosome 9 by analyzing the F2 population derived from crossing conventional and high-palmitic lines. Markers of these loci were located close to gene encoding 3-ketoacyl-ACP synthase II associated with high levels of palmitic acid (Pérez-Vich et al., 2016). A recent GWAS study performed in our lab further helped to identify potential genetic markers (SNPs) associated with 12 fatty acids (Chernova et al., 2021).

Genes controlling tocopherol composition were mapped in sunflower variants by analyzing experimental crosses (BC, F2). It was demonstrated that this trait is controlled by two recessive/dominant genes *Tph1* and *Tph2*, located on chromosomes 1 and 8 encoding 2-methyl-6-phytylbenzoquinol methyltransferase and γ -tocopherol methyltransferase explaining up to 97% of phenotypic variance (Tang, Hass, et al., 2006; Vera-Ruiz et al., 2006). However, several studies indicate that this trait, besides the two major effect genes, might include additional minor non-allelic genes located on chromosomes 4, 9, 14, and 16 (García-Moreno et al., 2012a; Hass et al., 2006). Thus, despite the suggested mono-/dihybridism inheritance of oleic acid content and tocopherol composition, additional loci were reported for both traits' indicating the importance of the genetic background for traits being mainly under mono-/oligo genic control (García-Moreno et al., 2012a; Hass et al., 2006).

As climate change remains a serious challenge for plant breeding, more studies focus on QTLs of drought and high salinity resistance. Fourteen QTLs that determine high salinity resistance were identified by analyzing experimental crosses of wild sunflowers (Lexer et al., 2003). In cultivated sunflower variants, six candidate loci were linked with salt tolerance: the loci encoding genes involved in the mineral ion uptake (Lai et al., 2005). Several studies were performed with an aim of detecting QTLs associated with the agronomically important traits under different water regimes by analyzing experimental crosses (Poormohammad Kiani et al., 2007, 2008, 2009). As a result, a QTL associated with yield stability across water-stressed and standard conditions was identified (Poormohammad Kiani et al., 2009). Additionally, water use efficiency was assessed under drought conditions, and respective QTLs were identified on chromosomes 13, 15, and 9 cumulatively explaining up to 41% of trait variance (Adiredjo et al., 2014). A recent GWAS study identified five genetic markers associated with the thousand-kernel-weight traits, an important economic parameter, under drought response (Gosseau et al., 2019).

In conclusion, sunflower genomics includes numerous successful efforts leading to the dissection of genetic features underlying agronomically important

traits. These traits include herbicide and pathogen resistance, hybrid breeding oil quality and climate adaptations. Such progress not only improved our understanding of the genetic control mechanisms of various agronomic traits but also led to the identification of the corresponding genetic markers. This in turn has formed a substantial basis for the development of the marker-assisted selection approaches for sunflower improvement.

2.5 Genetics of agronomically important rapeseed traits

Similar to sunflower, rapeseed is well studied in terms of genetic diversity as it is the second-largest oilseed crop in the world. The first genetic maps for this crop were constructed in the mid-1990s based on the RFLP and RAPD markers (Ferreira et al., 1994; Parkin et al., 1995; Uzunova et al., 1995). However, the rapeseed genome was first sequenced and assembled relatively recently (Chalhoub et al., 2014). As in the case of sunflower, this knowledge facilitates genetic mapping and the development of markers for agronomically important traits in rapeseed.

There are six CMS systems identified in rapeseed, of which *ogu* and *pol* carrying lines are the most popular ones used to produce hybrid seeds (Anisimova & Dubovskaya, 2020). Thus, fertility restorer genes, as well as corresponding markers, were developed to facilitate hybrid breeding. For *ogu* fertility restorer gene *Rfo* introduced from radish (*Raphanus sativus*) RAPD markers were developed (Delourme et al., 1994). Further, a fine-mapping placed this locus to chromosome 19 and allowed to identify of the candidate gene encoding pentatricopeptide repeat (PPR) protein family member (X. Hu et al., 2008). For *pol* CMS derived from cultivar Polima, the monogenic dominant restorer gene *Rfp* was mapped on chromosome 18, by using RFLP-, RAPD-based genetic maps (Jean et al., 1997). A recent study focusing on fine mapping of *Rfp* by analyzing a large NIL population revealed a candidate gene encoding the PPR family protein (Z. Liu et al., 2012).

For oil and oilcake quality improvement, the genetic associations found two traits that significantly reduce the oil quality, namely glucosinolate content and erucic

acid. QTL mapping of these traits allowed to identify QTLs located on chromosomes 9 (A09), 12 (C02), 17 (C07), and 19 (C09) were considered as major effect loci controlling erucic acid content (Howell et al., 2003; Uzunova et al., 1995; J. Zhao & Meng, 2003). Recent GWAS studies confirmed loci located on chromosomes 2 (A02), 9 (A09) and 12 (C02) explaining up to 40% of phenotypic variance (Harper et al., 2012; F. Li et al., 2014; D. Wei et al., 2019). The genes located in these regions included ones homologous to the *HAG1* of *Arabidopsis thaliana* encoding transcription factor involved in the glucosinolate biosynthesis regulation (Gigolashvili et al., 2007; F. Li et al., 2014). In addition to major effect loci, several minor effect loci carrying genes involved in glucosinolate metabolism have also been mapped (Gubaev et al., 2020; Kittipol et al., 2019; Y. Liu et al., 2020; D. Wei et al., 2019), thus making it possible the development of marker-assisted selection towards reduction of glucosinolate content. For erucic acid, QTL studies identified two loci controlling its content (Ecke et al., 1995; Jourdren et al., 1996; Thormann et al., n.d.). Recent GWAS studies demonstrated that *BnaA.FAE1* and *BnaC.FAE1* associated with the erucic acid content are located on chromosomes 8 (A08) and 13 (C03) and explain up to 25% of phenotypic variance (F. Li et al., 2014; C. Qu et al., 2017; D. Wei et al., 2019; Q. Zhu et al., 2019). These genes encode fatty acid elongases and were shown to be involved in the biosynthesis of erucic acid in rapeseed (N. Wang et al., 2008; G. Wu et al., 2008). Additional minor effect loci were also identified (Havlickova et al., 2018; F. Li et al., 2014; C. Qu et al., 2017; C.-M. Qu et al., 2015; D. Wei et al., 2019; Q. Zhu et al., 2019) indicating that such a complex trait should be controlled by the implementation of genomic selection approaches in the breeding program.

Several studies further focused on the mapping of other oil-related traits – fatty acid content and tocopherol composition. As in the case of the sunflower, the main direction of the selection effort focused on reaching high content of oleic acid and improvement of tocopherol composition. QTL mapping revealed a major effect locus controlling oleic acid content in rapeseed on chromosome 5 (A5), explaining up to

76% of the phenotypic variance. This locus was shown to carry a gene encoding fatty acid desaturase (X. Hu et al., 2006; Schierholt et al., 2000). More recent GWAS studies confirmed the role of this fatty acid desaturase in oleic acid level control (Gacek et al., 2017) and allowed to identify additional loci carrying genes encoding acyl carrier protein and 3-ketoacyl-CoA synthase (Guan et al., 2019; Q. Zhu et al., 2019). For tocopherol composition mapping, one of the first studies identified two loci controlling tocopherol content explaining 43% of the phenotypic variance. Similarly, for tocopherol composition (α -/ γ -tocopherol ratio), additional two loci were identified explaining 35% of phenotype variance (Marwede et al., 2005). Association mapping studies further identified candidate genes *VTE3* and *PDS1*, encoding MPBQ methyltransferase and hydroxyphenylpyruvate dioxygenase involved in the biosynthesis of tocopherols (Fritsche et al., 2012; X. Wang et al., 2012). Recent associative transcriptomic studies that facilitate identification of molecular markers linked to trait-controlling loci further supported the role of *VTE4* encoding γ -tocopherol methyltransferase and controlling alpha-/gamma-tocopherol ratio (Havlickova et al., 2018).

As the diseases significantly affect yield preservation and productivity, many attempts have been made towards resistance genes and loci associated with quantitative disease resistance (QDR) identification (Neik et al., 2017; Ton et al., 2020). For clubroot disease caused by soil-borne protist *Plasmodiopora brassicae* few R genes were identified in *B. napus* by means of analyzing mapping populations derived from crosses between susceptible and resistant lines (Diederichsen et al., 2006; Fredua-Agyeman & Rahman, 2016; Manzanares-Dauleux et al., 2000). Notably, it was demonstrated that the clubroot resistance controlled by R genes could be overcome by pathogens (LeBoldus et al., 2012) thus several studies focused on discovering QDR loci were made. A total of 19 QDR loci have been identified by means of QTL mapping in rapeseed, explaining up to 80 percent of phenotypic variance (Werner et al., 2007). Recent studies performed by means of association mapping in diversity panels allowed the identification of more than 30 additional

QDR loci (Hejna et al., 2019; L. Li et al., 2016). For parasitic fungus *Leptosphaeria maculans* causing blackleg disease at least seven R genes have been mapped on two chromosomes 7 (A7) and 10 (A10) by analyzing experimental crosses derived from R gene carriers (Ansan-Melayah et al., 1998; Balesdent et al., 2001; Rimmer, 2006). As the qualitative resistance is overcome by the pathogen several recent studies have been supporting the fact that the QDR loci increase the effect of the R genes (Brun et al., 2010; Delourme et al., 2014). Thus more recent studies are focused on the mapping of QDR loci. As a result, more than 20 QDR loci have been identified by analyzing genetically contrasting collections and contrasting DH lines of rapeseed (Fu et al., 2020; Huang et al., 2016; R. Raman et al., 2020). Unlike for the *L. maculans* and *P. brassicae*, identification of R genes for parasitic fungus *Sclerotinia sclerotiorum* causing soft rot remains challenging due to the medium heritability of the *B. napus* resistance. As a result several attempts have been made to find QDR loci through assessment of genetic contrasting collections, DH mapping populations and integrative analysis based on previous studies (J. Li et al., 2015; L. Wei et al., 2016; J. Wu et al., 2013, 2016). Identified markers explained up to 32% of phenotypic variance (J. Wu et al., 2013) candidate genes were involved in oxidative stress response, secondary metabolites production, hormonal pathways functioning and pathogen recognition (J. Li et al., 2015; L. Wei et al., 2016; J. Wu et al., 2013, 2016). A very recent GWAS study identified 123 loci associated with rapeseed resistance to *S. sclerotiorum*. Additionally, genomic prediction model was developed with a predictive ability of 64% which in turn could be used for genomic selection for *S. sclerotiorum* resistance in rapeseed (Roy et al., 2021).

Crop preservation during the vegetation period for rapeseed is determined by pod shattering and resistance to lodging. For pod-shattering only recent studies based on the analysis of diversity panels and artificial mapping populations helped to identify candidate loci. Pod shattering demonstrates high broad-sense heritability (J. Liu et al., 2016). As a result, a major effect locus carrying *psr1* gene explaining 47% of the phenotypic variance was mapped on chromosome A09 (Z. Hu et al., 2012).

Additional GWAS study identified 12 additional QTL that jointly explained approximately 57% of the genotypic variation of pod shattering (H. Raman et al., 2014). Although lodging leads to a significant yield loss of up to 46% (Kendall et al., 2017), studies dissecting resistance to lodging appeared recently. By analyzing the diversity panel by means of an associative transcriptomic approach three genomic regions (explaining up to 30% of phenotypic variance) associated with the stem strength were identified. Candidate genes located in these regions were associated with plant cell wall biosynthesis and regulation of hormonal pathways (C. N. Miller et al., 2018). Another GWAS study revealed five loci associated with lodging, however, the proportion of explained variance ranged from 8 to 10 percent (L. Wei et al., 2017).

Rapeseed is significantly affected by the non-optimal water and temperature regimes, including drought, salinity and cold, therefore the breeding for abiotic stress tolerance remains crucial under a rapidly changing climate (Ahmar et al., 2019). Thus several recent studies aimed at identifying genetic markers for abiotic stress resistance have been made. A DH mapping population was used to map waterlogging and drought tolerance. As a result, 26 QTL and 31 QTL were identified explaining up to 23 % of phenotypic variance (Z. Li et al., 2014). Two studies based on GWAS identified novel loci and genes that determine drought and salt tolerance. Twenty-five QTLs explained up to 9.23% of salt tolerance and carried genes encoding aquaporins acting as water channels, transcription factors involved in abiotic stress response, proline biosynthesis enzymes (Wan et al., 2017). For drought stress tolerance two loci located on chromosomes 10 (A10) and 19 (C9) were identified; four genes encoding PPD5 regulating ROS levels and stomata closure as well as ribosomal protein (RP), multidrug and toxic compound extrusion (MATE) protein, and a histone superfamily protein previously shown to regulate abiotic stress responses were suggested as candidate ones controlling the trait (Shahzad et al., 2021).

Summarizing the information reviewed in the present chapter it should be concluded that the recent progress in genotyping and data analysis approaches

facilitated the dissection of the agronomically important traits in rapeseed. Significant progress was made both for monogenic traits, such as qualitative disease resistance and ability to restore fertility, and for polygenic traits including fatty acid composition, glucosinolate content and resistance to unfavorable environmental conditions. Identification of trait-associated genomic loci, in turn, facilitated the understanding of the genetic control of the traits including the identification of novel candidate genes. More practically, these studies revealed prospective genetic markers for marker-assisted and genomic selection in rapeseed.

2.6 Russian trends in rapeseed and sunflower breeding

Sunflower and rapeseed sowing areas rank first and third, respectively, among oilseed crops in Russia. It should be noted that for rapeseed and sunflower, up to 53 and 73 percent of planting material currently used in Russia is of foreign origin, which is explained by the insufficient development of local breeding companies and institutions. The leading Russian institute for sunflower breeding and the major governmental originator is VNIIMK (Pustovoit All-Russia Research Institute of Oil Crops). Besides VNIIMK institute that produces novel varieties of sunflower is Altai Research Institute of Agriculture and the Federal Center of Agriculture Research of the Southeast Region. Additionally for sunflower, private breeding companies including "Agroplasma", "Galaktika" and "Triumph" made a substantial contribution to new varieties' development. For rapeseed, the leading variety breeder is the All-Russian research institute of rapeseed, in addition VNIIMK and Siberian Federal Scientific Center for Agrobiotechnologies. Unlike sunflower, rapeseed breeding in Russia is currently supported by a single private breeding company "Astra" (Ivanov et al., 2020).

For sunflower, there are several breeding trends currently addressed in Russia. One of the top problems that face sunflower breeders is related to the resistance to the broomrape. Active attempts are made for pyramidisation of the resistance from race A to recent race G (Demurin et al., 2020; Ryzhenko et al., 2021). In the area of hybrid

breeding that requires herbicide resistance and involvement of CMS/Rf systems significant progress was made in the identification and introduction of herbicide resistance and Rf genes (Anisimova et al., 2021; Chelyustnikova et al., 2017; Demurin, Borisenko, et al., 2006).

Successful attempts were made toward the development of hybrids with improved oil quality, basically the high-oleic sunflowers with altered tocopherol composition (Demurin, Borisenko, et al., 2006; Demurin et al., 2016). Another important direction in sunflower breeding is the creation of the confectionary sunflower which is generally between edible and oil forms. It should be noted that confectionary sunflower breeding is mostly based on the populational varieties (Bochkovoy et al., 2020).

For rapeseed, the main breeding targets in Russia are oil quality, resistance to herbicides, yield preservation, and productivity. For oil quality, the low erucic and low glucosinolate varieties are developing in the VNIIMK institute. Additionally, chemical mutagenesis was applied to introduce a high oleic trait in Russian germplasm (Gorlova et al., 2020). It also should be noted that most of the Russian rapeseed registered varieties are represented by inbred lines, while foreign variety producers mostly rely on hybrid breeding (Anisimova & Dubovskaya, 2020). Resistance to lodging and pod shattering also remains important traits in Russia and are evaluated for new varieties of high interest for breeding (Bochkaryova, Strelnikov, et al., 2021). Further, VNIIMK recently started to initiate hybrid rapeseed breeding since the productivity of hybrids is higher compared to inbred lines (Bochkaryova et al., 2020). The first hybrid based on the *ogu* CMS system "Debut" was registered in 2020. Notably, the productivity of the hybrid was equaled to the ones obtained in the EU and significantly higher than the "Loris" standard variety (Bochkaryova, Gorlova, et al., 2021).

Despite the huge work on the development of the varieties that meet the industry's requirements, marker-assisted breeding is poorly used in scientific institutions and private breeding companies in Russia. Only a few studies have been

made for mapping agronomically important traits in rapeseed and sunflower in Russia (Usatov et al., 2014). Notably, a significant part of these studies was made in collaboration with Skoltech (Chernova et al., 2021; Goryunov et al., 2019; Goryunova et al., 2019; Gubaev et al., 2020, 2022). The present work is a part of an effort of introducing marker-assisted selection in Russia for rapeseed and sunflower improvement. This work is focused on the mapping of the seed glucosinolate content in rapeseed, tocopherol composition and oleic acid content as well as seed-related traits in sunflower.

Chapter 3. Materials and methods

3.1 Plant material

3.1.1 Rapeseed diversity panel

Rapeseed diversity panel consisted of 90 accessions from the All-Russian Research Institute of Oil Crops (VNIIMK) and included 43 winter and 47 spring inbred lines (Gubaev et al., 2020). The phenotype data on glucosinolate content was collected during three vegetation periods (2015–2016, 2016–2017, 2018–2019) at VNIIMK experimental station, GPS coordinates were 45°10'51" N and 39°02'42" E. Seeds of plants used for genotyping were collected during 2018–2019 vegetation period.

3.1.2 Experimental crosses for oil-related traits mapping in sunflower

The plant material used to map tocopherol composition and oleic acid content consisted of genetically contrasting mutant lines VK195 and VK876 as well as wild type VK303 and VK101. According to hybridological analysis lines VK195 and VK876, carry mutant alleles *tph1*, *tph2* and *Ol* and thus express γ - and δ - tocopherol phenotype (~50% of γ - and ~50% δ -tocopherols) as well as demonstrate high oleic phenotype (>82% of oleic acid). Lines VK303 and VK101 carry wild type *Tph1*, *Tph2* and *ol* and demonstrate a high α - tocopherol phenotype (100% of α -tocopherol) as well as moderate content of oleic acid (30 - 45%). All lines were developed at the VNIIMK institute. Elite lines VK101 and VK303 are used to produce a simple interlinear middle-early sunflower hybrid "Typhoon" registered at VNIIMK (Trembak et al., 2018). While Lines VK876 and VK195 are used to produce an "Oxy" hybrid that synthesizes oil with increased oxidative stability (Demurin et al., 2016).

Lines VK195 and VK876 were used as female lines after hand emasculatation. Lines VK101 and VK303 were used as a source of pollen. The crosses were performed in the field (Krasnodar) in 2015. Individual F1 plants were selfed using

isolators to produce the F2 progeny (F2 seeds) in the field (Krasnodar) in 2018. Random inflorescence was selected from a plot. For subsequent phenotyping and genotyping procedures F2 seeds were randomly sampled per F1 sunflower head. In total 142 F2 seeds for cross VK195xVK303 and 144 F2 seeds for the cross VK876xVK101 were collected. Seeds were harvested at the physiological maturity stage (R-9) according to the sunflower growth stages classification (Schneiter & Miller, 1981). For phenotyping, a half-seed technique was applied (Y. N. Demurin et al., 2004; Vera-Ruiz et al., 2005). In particular, a single seed was cut in half with a razor, and the first part containing the embryo was left for germination for subsequent genotyping, while the second part was used for phenotyping.

3.1.3 Sunflower diversity panel

A panel of 601 sunflower accession was provided by N.I. Vavilov Institute of Plant Genetic Resources (VIR, St. Petersburg, Russia) - 255 accessions, V.S. Pustovoit All-Russia Research Institute of Oilseed Crops (VNIIMK) - 199 accessions, and LLC Agroplasma Seed and Breeding Company (Krasnodar, Russia) - 147 accessions. All plants were grown and seeds were collected in the Krasnodar region in Russia.

All accession were diploid, $2n = 34$, seeds were stored for 1–3 years. Plants were grown in fields of leached black earth soil type. The sunflower was sown following the preceding crop, fall wheat, at the seeding rate of 40,000 plants per hectare. Sowing was carried out according to the following sowing system: 70×35 cm, a single plant per planting pit. Farming techniques, as commonly used for sunflower. Each line was grown on the plot with an area of 9.1 m^2 .

3.2 Phenotyping procedures

3.2.1 Glucosinolate measurement

For glucosinolate measurement, a modified colorimetric analysis was applied (Thies, 1982). For each measurement, 5 g of seeds were collected from three plants of

each line from a single plot during each of the three years. Seeds were homogenized using a mechanical grinder, then 0.2 g of the homogenate was used for extraction according to the previously published protocol (Gubaev et al., 2020). The obtained extract was subjected to measurement of absorbance at 420 nm on a spectrophotometer. The concentration was calculated based on the calibration curve obtained using the standard samples with known concentrations.

3.2.2 Tocopherol composition and oleic acid content measurement

To determine tocopherol composition, a thin-layer chromatography (TLC) with subsequent densitometry was applied. Extraction of the tocopherols was made according to the previously described protocol (Demurin, Efimenko, et al., 2006; Y. N. Demurin et al., 2004). For the detection of tocopherols, the plate was sprayed with a freshly prepared mixture of 0.1% ferric chloride (FeCl_3) and 0.25% α, α' -dipyridyl in absolute ethanol (Emmery-Engel's reagent), taken by volume in a ratio of 1:1. The stained plates were scanned and the relative composition in tocopherol forms was quantified by measuring the relative intensity of the color of tocopherol forms (in %) using a Sorbfil videodensitometer (software version 1.5.0).

To determine the fatty acid composition of the oil and relative content of oleic acid, a half seed was ground in a coffee grinder, thoroughly mixed, and a 0.5 g sample was taken for extraction in 4 ml of hexane. Then 2 ml of miscella were taken for methylation according to the standard method (Demurin, Efimenko, et al., 2006; Y. N. Demurin et al., 2004). Chromatographic analysis was performed on a Khromatek-Kristall 5000 gas chromatograph with an automatic dispenser DAZH-2M on a SolGelWax 30 m \times 0.25 mm \times 0.5 μm capillary column in a flow of helium carrier gas, at a speed of 25 cm/s, with temperature programming in within 185-230 $^\circ\text{C}$. The preparation of methyl esters and their chromatography and subsequent data analysis were performed in accordance with standard methods (Demurin, Efimenko, et al., 2006; Y. N. Demurin et al., 2004).

3.2.3 Seed-related traits assessment

To estimate the seed and husk area, seeds were placed on the plate. For each line, 10 seeds were analyzed per plate. The next plate was scanned by means of PRDU-02 mobile X-ray diagnostic installation jointly developed by Agrophysical Research Institute and St. Petersburg State Electrotechnical University LETI (Arkhipov et al., 2019). Image analysis was performed by SeedRentgen software (Брач et al., 2015). The mean values of husk and seed area were calculated for each of the plates (accession), given the area of one pixel of each photo corresponded to $100\ \mu\text{m} \times 100\ \mu\text{m}$ ($0.1\ \text{mm}^2$) of the plate area. Next, a ratio between seed area and husk area was calculated based on the rounded values for subsequent analysis. To assess the effect of the collection on the husk and seed area as well as seed to husk ratio analysis of variance was performed. To assess the correlation coefficient between husk and seed area Pearson correlation was estimated. Mann–Whitney U test was performed to assess the difference in seed area and husk area and seed to husk ratio between collections.

3.3 Genotyping procedures and SNP calling

3.3.1 Rapeseed genotyping and SNP calling

For DNA extraction 1–2 leaves of seedlings germinated within two weeks under room conditions were used. Three independent plants of each line were used for genotyping procedures. DNA was extracted according to the CTAB protocol using the NucleoSpin Plant II plant DNA extraction kit (Macherey-Nagel, Dylan, Germany). Qubit 3.0 Fluorometer (Thermo Fisher Scientific, Waltham, MA, USA) and gel electrophoresis were used to estimate the quality and concentration of the DNA. Extracted DNA was used to prepare the genotyping-by-sequencing (GBS) library as previously described (Chernova et al., 2021; Goryunov et al., 2019). PstI and MspI restriction enzymes were used to cut DNA. Sequencing was performed in the HiSeq 4000 (Illumina, San Diego, CA, USA). Raw sequence data are available on NCBI SRA under the project number PRJNA645178.

To find SNPs raw reads were first demultiplexed according to the barcodes which identify sample ID using Axe-demultiplexer software version 0.3.3 (Murray & Borevitz, 2018). Trimmomatic software version 0.36 was used to perform read filtering according to the following parameters: low-quality bases (Phred score < 20) as well as sequences corresponding to Illumina adapters and reads of less than 40 bp were removed. Filtered reads were mapped on the most recent version of the reference genome of *Brassica napus* cultivar 'ZS11' (F. Sun et al., 2017) using Burrows-Wheeler aligner (H. Li & Durbin, 2009) version 0.7.17 with default parameters. For SNP calling, a modified version of the pipeline (https://github.com/RimGubaev/GATK_pipeline_customized) based on Genome Analysis Toolkit software version 4.1 (McKenna et al., 2010) was applied. The biallelic SNPs that passed the following filters were used for subsequent analysis: Minor allele frequency (MAF) > 0.03 and maximum missing data of 30%, sequencing depth was of at least 10 reads per position. Additionally, SNPs were filtered by maximum heterozygosity of 40%, as it was previously discussed fact of the high amount of heterozygous positions leads to read misalignment due to the amphidiploid nature of the rapeseed genome which results in the presence of highly homologous regions (Malmberg et al., 2018).

3.3.2 Sunflower genotyping and SNP calling

To perform genotyping of the sunflower plants, the half-seeds left after cotyledon separation were germinated in rolls of filter paper. DNA was extracted from the cotyledon leaves using the NucleoSpin Plant II kit (Macherey-Nagel) according to the manufacturer's protocol. Modified protocol (Elshire et al., 2011) was used to prepare a genotyping-by-sequencing (GBS) library as previously described (Chernova et al., 2021; Goryunov et al., 2019). Illumina HiSeq 4000 was used to sequence prepared libraries of single-end reads with a length of 150bp. For F2 crosses, parental plants were genotyped in at least seven biological replicates. Raw

sequence data for F2 cross and parental lines are available on NCBI SRA under the project number PRJNA742188.

For SNP calling TASSEL-GBS pipeline v2 was used. Read mapping was performed using bowtie2 (Langmead & Salzberg, 2012) aligner with `-very-sensitive` parameter. The most recent assembly XRQ2.0 (<https://www.heliagene.org/HanXRQr2.0-SUNRISE/>) of the sunflower genome was used as a reference (Badouin et al., 2017) genome. Prior to mapping, reads derived from each parental line's replicate were merged. Next for genetic map construction only SNPs that were homozygous within each parent and polymorphic between the parents were selected. SNPs possessing less than 10% or more than 90% of heterozygosity level as well as ones that were genotyped in less than 20% of accessions, were discarded before the imputation procedure as previously recommended (Fragoso et al., 2016).

To perform SNP calling for diversity collection of the sunflower, a modified GATK pipeline was applied. Variants were filtered using hard filtering parameters: Map quality (MQ) > 36, QD > 24, and MQRankSum < 2, ensuring that reads were mapped to a unique place in the reference, that the reads carrying both alleles were comparable in terms of mapping quality (MQRankSum), and that the actual variants were called with high quality (QD). To select SNPs for population structure and association mapping analyses of seed-related traits missing calls rate < 0.3, variant depth (DP) > 4 and minor allele frequency (MAF) > 0.01 were applied, resulting in 15,068 SNPs.

3.4 Population structure analysis

To identify the number of subpopulations in sunflower and rapeseed datasets ADMIXTURE software version 1.3 (D. H. Alexander et al., 2009) was applied with the number of clusters ranging from 1 to 10. The PLINK software version 1.9 (Purcell et al., 2007) was used to perform principal component analysis with standard parameters, the number of principal components was set to 10, PCs were estimated

using the variance-standardized genetic relationship matrix. To estimate linkage disequilibrium r^2 was calculated among SNP pairs located within a 1500 kb window using PLINK software version 1.9.

For diversity comparative analysis of rapeseed WGS data for 54 rapeseed accession from European, Chinese, Australian, Japanese, Moroccan, and Indian collections (Malmberg et al., 2018). Lines were of different ecotypes (winter, semi-winter, and spring), SNP calling of raw WGS data was performed as described above (see chapter 3.3.1). To compare the genetic diversity of the sunflower, a raw vcf file including 1065 wild, 20 landraces, and 289 cultivated accessions (Hübner et al., 2019). Bcftools software version 1.9 (H. Li, 2011) was applied to find common SNPs among the datasets between the studied cohorts and the international accessions of rapeseed and sunflower.

3.4 Association mapping and candidate gene identification

3.4.1 Association mapping of glucosinolate content

Compressed mixed linear model (cMLM) implemented in TASSEL software version 5.2 (Bradbury et al., 2007) were used to find genetic associations to glucosinolate content. Kinship matrix calculated with centered IBS (Identity by state) method as well as first five principal components were included in the model to account for kin relationships and population structure, respectively. Phenotype data for each year were analyzed independently. To find the most reproducible SNPs across three years, the sum of the p-values was substituted by the corresponding densities of the Irwin-Hall distributions taking into account that the sum of p-values was formed by three uniform distributions. This was done using Unifed R package's Dirwin.hall function (Quijano Xacur, 2019). The adjusted p-value with Bonferroni correction was set to 0.0000041 (0.05/12,226, where 12,226 is the number of SNPs) to consider associations. An additional softer threshold was set to 0.0005. The scanning for potential candidate genes was performed using SnpEff software version 4.3 (Cingolani et al., 2012). A frame of 100 kb upstream and downstream of the

significant SNPs was used, as such a threshold was previously used for scanning candidate genes in rapeseed (Guan et al., 2019; Q. Zhou et al., 2018; Q. Zhu et al., 2019).

To compare previously identified loci with ones found in the present study an information on the exact physical location of the SNPs associated with glucosinolate content was collected from previously published data (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020; C.-M. Qu et al., 2015; D. Wei et al., 2019). Next for each of the SNPs identified in the present study, corresponding positions were identified in the *Brassica napus*'s Darmor bzh assembly (Chalhoub et al., 2014) as it was used in the previous studies. To do so the 50bp upstream and downstream flanking sequences of the SNPs were extracted and mapped to the Darmor bzh assembly version 4.1 using Blast software version 2.2.18 (Camacho et al., 2009). The following filters were used to identify the SNPs common for two assemblies: e-value $< 1 \times 10^{-40}$. alignment length > 90 . The frame of 100 kb was used to find the genetic regions in common with the previous studies.

3.4.2 Association mapping of the seed morphology traits in sunflower

To perform association mapping of the seed-related traits a compressed mixed linear model (cMLM) was implemented in TASSEL software version 5.2 (Bradbury et al., 2007). Kinship matrix calculated with centered IBS (Identity by state) method as well as first five principal components were included in the model to account for kin relationships and population structure, respectively. Additionally, the collection of origin was added to the model as a covariate as it was demonstrated that it is a factor significantly affecting seed and husk area as well as seed to husk ratio (ANOVA p-value $\ll 0.05$). The adjusted p-value with Bonferroni correction was set to 0.0000033 (0.05/15,058, where 15,058 is the number of SNPs) to consider associations. An additional softer threshold was set to 0.0001. The scanning for potential candidate genes was performed using SnpEff software version 4.3 (Cingolani et al., 2012). A frame of 100 kb upstream and downstream of the significant SNPs was used as on average SNP pairs located within this frame

demonstrated a high level ($r^2 > 0.5$) of linkage (for details see Chapter 6.3, Figure 6.3.1).

3.5 Genetic map construction

Before genetic map construction, missing genotypes were imputed with default parameters using the LB-impute pipeline (Fragoso et al., 2016). Imputed SNP genotypes were encoded as the A, B, and H, corresponding to maternal, paternal and heterozygous genotypes, respectively using a custom R script (https://github.com/RimGubaev/vcf_to_qtl). SNPs genotyped in less than 80% of individuals as well as ones with a significant segregation distortion (Bonferroni-corrected χ^2 goodness-of-fit test P-value < 0.05) were discarded. Additionally, Individuals genotyped by less than 80% of SNPs were removed. To construct a genetic map R/qtl software (Broman et al., 2003) was applied. Data on physical chromosomes available for SNPs was used to preorder the loci within chromosomes. Kosambi mapping function was implemented to order markers within chromosomes. Additional filtering for SNPs within each chromosome was performed using the `droponemarker()` function; markers with LOD scores equal to or above -20 were discarded. To estimate the concordance between physical and genetic marker order the Pearson correlation coefficient and the corresponding significance level were calculated using `cor()` and `cor.test()` functions, respectively, in the R base package between physical and genetic distances.

3.6 QTL mapping of tocopherol composition and oleic acid content

Quantitative and qualitative approaches were used to map genetic markers of tocopherol composition and oleic acid content. For quantitative mapping of the proportion of each of the tocopherol classes and oleic acid content were considered as independent observations. Analysis of variance (ANOVA) implemented in R's base statistical package was used to calculate the proportion of variance explained by genetic markers. For these traits, the interval mapping approach for non-normally

distributed traits based on Haley–Knott regression was applied using the `scanone()` function from `r/qtl` package. Additionally, composite interval mapping was used to map oleic acid content. For qualitative mapping of *Tph1* loci the phenotypes of plants that belong to the α (*Tph1*/_; *Tph2*/_) and γ (*Tph1*/_; *tph2/tph2*) classes were set to 1, while the α/β (*tph1/tph1*; *Tph2*/_) and γ/δ (*tph1/tph1*; *tph2/tph2*) classes were set to 0. For qualitative mapping of the *Tph2* the phenotypes of plants that belong to the α (*Tph1*/_; *Tph2*/_) and α/β (*tph1/tph1*; *Tph2*/_) classes were set to 1, while the β (*Tph1*/_; *tph2/tph2*) and γ/δ (*tph1/tph1*; *tph2/tph2*) classes were set to 0.

The phenotypes of plants that did not belong to any of the classes based on the tocopherol composition were set to NA. For qualitative mapping of the *Ol* gene phenotypes of plants that demonstrated oleic acid content equal to or higher than 83% were set to 1 while those that demonstrated less than 83% were set to 0. To scan for loci associated with *Tph1*, *Tph2* and *Ol*, the interval mapping approach for binary traits based on Haley–Knott regression implemented in the `r/qtl`'s `scanone()` function was applied. Permutation analysis for the logarithm of odds (LOD) carried with 1,000 iterations was performed to identify a significance threshold corresponding to the 99 percentile of the permuted LOD values. To calculate 1.5-LOD confidence intervals for most significant markers `lodint()` function in `r/qtl` was applied.

3.7 Verification of the genetic markers for tocopherol composition and oleic acid content

To validate genotypes of polymorphic SNPs obtained with GBS, significant SNPs associated with *Tph1*- and *Tph2*-determined phenotypes were selected. For primer design, 500bp upstream and downstream sequences were extracted for SNPs within 1.5-LOD interval bedtools v2.27.1 (Quinlan & Hall, 2010). Primer3 plus (Untergasser et al., 2012) was used to design primers for amplification of the regions and subsequent Sanger sequencing. To test primers for dimer formation and self-complementarity OligoCalc tool was applied (Kibbe, 2007). PrimerBlast software (Ye et al., 2012) implemented in the PlantEnsemble genome browser was

used to test the specificity of the primers within sunflower genome version XRQ2.0. Two criteria were used to select markers, the first is the high LOD scores within a 1.5-LOD confidence interval, second is the ability to construct unique primer pairs matching unique regions. As a result, four pairs for amplification were constructed (Table 3.7.1).

Table 3.7.1 – Primers used to amplify loci carrying significant SNPs associated with *Tph1*- and *Tph2*-associated genotypes.

Primer name	Sequence 5'-3'	Tm used for amplification	Product Size, bp
S1_55196434_Tph1_VK195xVK303_F	ACATGGTTTCTTATCATTTGCAC	59.0 C	420
S1_55196434_Tph1_VK195xVK303_R	ACCGGATATTTGACAAAGTGC		
S8_30578572_Tph2_VK195xVK303_F	TGACTTACTTGGTCGAGCCG	60.5 C	416
S8_30578572_Tph2_VK195xVK303_R	GCACGTACCCGATTCCTTG		
S1_71748138_Tph1_VK876xVK101_F	ATCCTCCACAACCCAACACG	60.0 C	459
S1_71748138_Tph1_VK876xVK101_R	GAAAGCATACTTTGGGCGACT		
S8_23941299_Tph2_VK876xVK101_F	TCTCGGATTACAGTGGTTCGA	59.5 C	287
S8_23941299_Tph2_VK876xVK101_R	GAAAACGATGGGGTTCTGG		

To validate markers 34 F2 plants from the mapping population eight plants for each of the following groups were selected: those carrying the paternal genotype, carrying the maternal genotype and plants carrying heterozygous genotype according to GBS data. Five parental plants were genotyped to validate paternal and maternal genotypes. Direct Sanger sequencing of the amplicons was performed, and Ugene software (Okonechnikov et al., 2012) was applied to analyze sequencing results. To test the concordance of the genotypes obtained by two different methods, we

compared the GBS-derived genotypes of the markers with the ones obtained by Sanger sequencing and estimated the proportion of the matching genotypes.

In addition to the SNP variant validation, the ability to predict *Ol*-, *Tph1*- and *Tph2*-associated phenotypes was tested. To do so independent sample of 20 lines was genotyped. Among them, eight were high oleic (>83%), and 12 represented wild type, regarding tocopherol composition all four classes were represented by five lines. Each line was sequenced in five replicates. SNP calling was performed jointly with each of the crosses in order to capture the genetic polymorphisms common between the test sample and each of the crosses. SNP calling was performed as described in chapter 3.3.2. Next, 10 most significant markers were selected for each trait/cross combination based on the QTL-mapping results. For each marker, a proportion of the genotypes in the test sample was calculated as well as the proportion of the correctly predicted phenotypes associated with *Ol*, *Tph1* and *Tph2* was estimated.

Chapter 4. Genetic characterization of Russian rapeseed collection and association mapping of glucosinolate content

4.1 Introduction

In the present part of the study, a genotyping of the rapeseed collection from the VNIIMK was performed with the aim of genetic characterization of the collection and the mapping of the agronomically important trait – glucosinolate content. To do so 90 genetically and phenotypically contrasted rapeseed accessions consisting of 47 spring and 43 winter accession were analyzed.

Glucosinolates represent secondary metabolites that on the one hand act as part of the defense against plant pathogens and on the other hand negatively affect the higher animals when consumed as a part of the diet, additionally glucosinolates contribute to the bitter taste of Brassicas. Thus the reduction of the glucosinolates is one of the key breeding targets in rapeseed production (Walker & Booth, 2001; Wieczorek et al., 2018). As it was the first study that included the genotype data of the diversity panel of Russian rapeseed, a population structure was described for the studied cohort. Additionally, we compared the genetic diversity of the studied cohort with the available whole-genome sequencing data available at the end of 2018 (Malmberg et al., 2018). The phenotyping procedures were performed during the three vegetation periods (2015-2016, 2016-2017, 2018). The collected data was used for the association mapping and identification of loci responsible for the glucosinolate content as well as for scanning for the potential candidate genes. Additionally, we performed the comparison of the identified loci with previously identified ones; this comparison showed that the loci identified in the present study are new. The results presented in the following chapter were published in the *Genes* journal (Gubaev et al., 2020).

4.2 Genetic characterization of VNIIMK rapeseed collection

To describe the genetic diversity of the studied cohort, the GBS approach was applied. As a result, we identified 160257 raw biallelic SNPs. After the filtering procedures, 12226 SNPs were left for the subsequent analysis, which is approximately 7-8 times lower than it was previously made by using similar reduced representation sequencing approaches (Gacek et al., 2017; Lees et al., 2016). Which on the one hand is explained by the relatively strict filters used to subset the genotypes, and on the other could be the result of a relatively small sample size capturing a smaller amount of genetic diversity.

Collected genotype data was used to estimate the number of the potential subpopulations, to do so the ADMIXTURE algorithm was applied (Figure 4.2.1). A significant drop in the cross-validation error indicating the number of potential subpopulations was observed while dividing a studied cohort into two, three, five and seven clusters (K) (Figure 4.2.1 A).

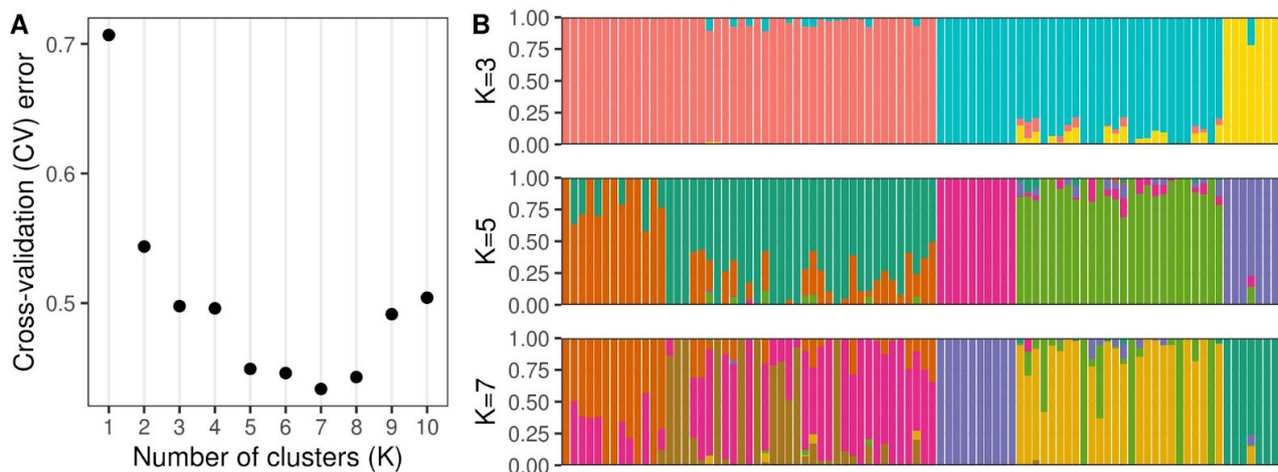


Figure 4.2.1 – Results of the population clustering assessment by the ADMIXTURE software. (A) Cross-validation error value for K ranging from 1 to 10. (B) ADMIXTURE bar plots correspond to subpopulations at K = 3, 5, 7, each bar corresponds to the rapeseed line; colors indicate the genetic admixtures for each of the studied lines. Red, blue and yellow bars reflect genetic admixture specific to spring, winter and yellow-seeded winter rapeseed accessions at K = 3.

At $K = 2$ the clusters were represented by the winter and spring ecotypes. A significant genetic difference between the spring and winter rapeseed accessions was expected and previously described as the major factor explaining the genetic structure since these two ecotypes demonstrate different growth habits as the winter rapeseed is sown in late autumn and requires vernalization while the spring ecotype is sown in the early spring (Gazave et al., 2016). Such segregation is explained by selection which led to the limiting of the rapeseed diversity and resulted in the separation of these ecotypes (Gazave et al., 2016; D. Wei et al., 2017; D. Wu et al., 2019). The principal component analysis confirmed this observation as the first principal component explaining 34.7% of genotype variance and also divided the studied sample into winter and spring ecotypes (Figure 4.2.2 A).

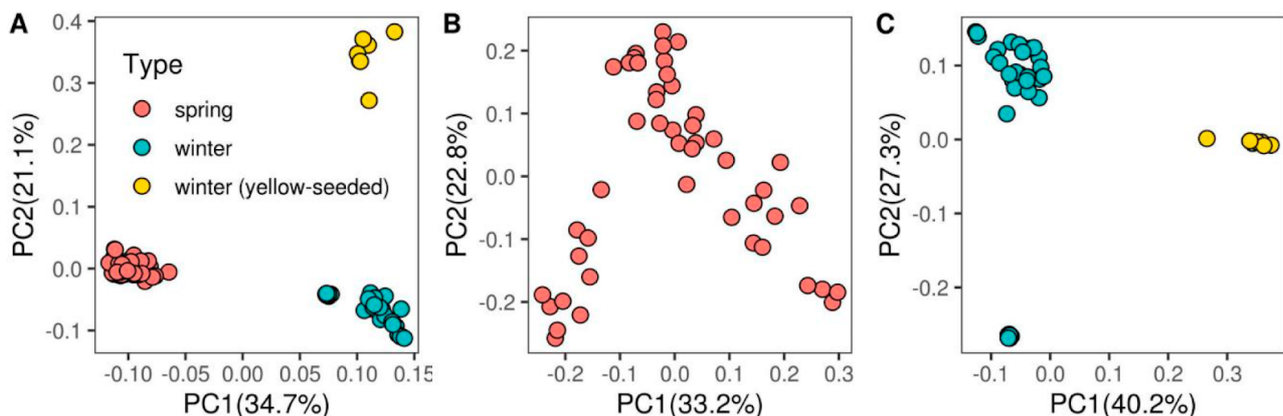


Figure 4.2.2 – Principal component analysis of the population structure of Russian rapeseed lines. PCA analysis was performed for the whole cohort (A) and separately for spring (B), and winter (C). Red, blue and yellow colors correspond to spring, winter and yellow-seeded winter rapeseed accessions, respectively.

A second significant drop in the cross-validation error was observed at $K = 3$; such segregation allowed additional distinguishing between yellow- and dark seeded winter lines (Figure 4.2.2 B). The principal component analysis also distinguished these groups by the second principal component explaining 21.1% genotype variance. Such segregation by the seed color was quite a new observation, as the color of the

seed coat is controlled only by several loci in *Brassicas* (HAWK, 1982; H. Zhao et al., 2019). This presumably could be explained by the recent breeding history of yellow-seeded accessions in VNIIMK.

Although the substantial drop in CV was also observed at $K = 5$ and 7 , such segregation did not correspond to any biochemical or phenotype data of VNIIMK breeders. The principal component analysis also did not reveal any clear clustering while analyzing the spring accessions separately (Figure 4.2.2). A separate principal component analysis of the winter accession revealed three clusters. The first PC explaining 40.2% of genotype variance is expectedly distinguished between yellow- and dark-seeded accessions. While the second PC explaining 27.3% of genotype variance additionally divided dark-seeded winter rapeseeds into two subpopulations, there was no explanation for this according to the breeders' data.

In the next step, we estimated the linkage disequilibrium (LD) decay based on the squared correlation coefficient (r^2) calculated among SNPs located within 1500kb windows for the whole genome (Figure 4.2.3 A). The proportion of SNP pairs with the r^2 value greater than 0.25 within 30 kb bins was also estimated (Figure 4.2.3 B). Additionally, LD was calculated separately for the A subgenome derived from a turnip (*B. rapa*) and cabbage (*B. oleracea*) (Figure 4.2.3 C).

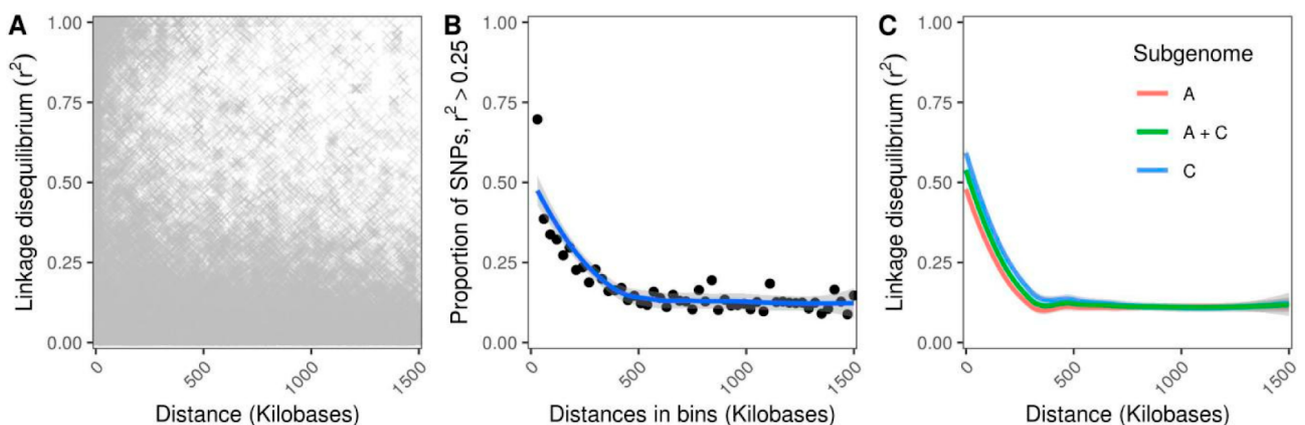


Figure 4.2.3 – Linkage disequilibrium (LD) decay in the studied rapeseed lines. (A) LD decay across genetic marker (SNP) pairs. Each cross corresponds to the r^2 value between a pair of SNPs within 1500kb. (B) The proportion of SNP pairs with $r^2 > 0.25$ was calculated for the whole genome. Each dot indicates the proportion of SNP

pairs in the 30 kb bin. (C) LD decay for A and C subgenomes. Colored lines on panels B and C represent loess curves. Gray markers correspond to the 95% confidence interval.

For the entire genome at $r^2 = 0.25$ size of the LD blocks equaled 218.8 ± 14.5 kilobases (kb). Our findings demonstrated that at linkage cutoff of $r^2 = 0.25$ LD decay was longer for the C subgenome and equaled 265.1 ± 13.4 kb and shorter for the A subgenome equalling 177.1 ± 19.5 kb. Such difference in the LD decay between A and C subgenomes was repeatedly reported for the collections that did not include a substantial amount of accessions from Russian gene banks (Qian et al., 2014; D. Wei et al., 2017; M. Zheng et al., 2017). Such differences were explained by the longer chromosomes of the C subgenome compared to ones from the A subgenomes (F. Sun et al., 2017).

As the present study was the first to include a significant amount of the rapeseed accession from a Russian gene bank genotyped in a high-throughput manner we performed a comparison of the genetic diversity of studied accessions with international ones from other collections for which WGS data was available at the end of 2018 (Malmberg et al., 2018). As a result 54 geographically and ecotypically diverse rapeseed lines were selected for the comparison. For these 54 accessions a WGS data was subjected to the same pipeline used for VNIIMK collection, resulting in 4,037,572 unfiltered SNPs which were joined with raw SNP data for VNIIMK collection. After filtering this joint dataset by the sequencing depth and missing values, common 20,848 SNPs polymorphic in at least one dataset were used to perform PCA analysis. The first and second principal components that jointly explained 38.7 % of genotype variance demonstrated that the international lines were clustered based on the collection of origin as well as the ecotype (Figure 4.2.4 A). This is in agreement with the previously published results where the collection of origin and ecotype served as the major factors affecting population structure in

rapeseed (Bus et al., 2011; Gazave et al., 2016; D. Wei et al., 2017; D. Wu et al., 2019).

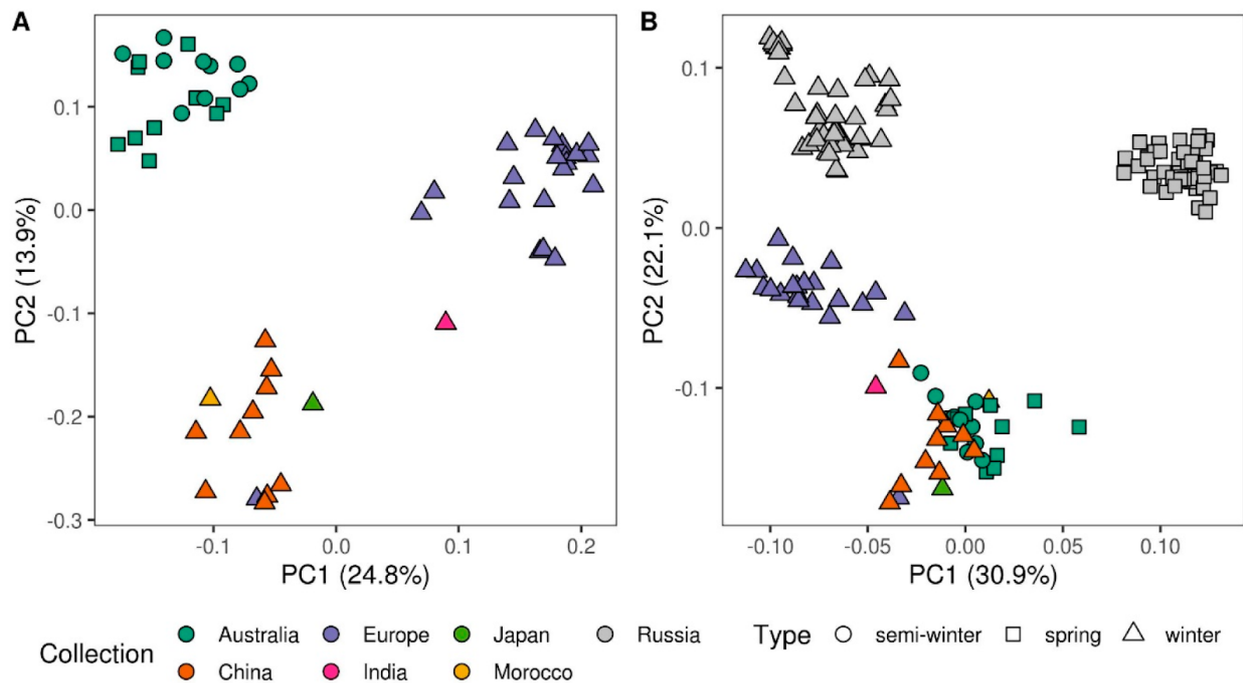


Figure 4.2.4 – Principal component analysis (PCA) plots reflecting the population structure of international and Russian collections. (A) The population structure of international lines is used in this study. (B) The comparison of the population structure of international and Russian lines. Shape corresponds to the ecotype, and colors represent the collection.

PCA analysis of the joint sample demonstrated the separation of the winter, spring and semi-winter accession by the first PC explaining 30.9% of genetic variance. The second PC explaining 22.1% genetic variance revealed clear differences between VNIIMK accessions and international ones (Figure 4.2.4 B). This in turn indicates a great potential for increasing the genetic diversity of collection for breeders and to find new genetic determinants for agronomically important traits. Additionally, this information could be used further for a deeper investigation of the rapeseed breeding history.

4.3 Measurement of the glucosinolate content

To perform the measurement of the glucosinolate, a modified chloropalade method followed by colorimetric analysis was applied at VNIIMK. The information on the glucosinolate content was collected for each of the 90 accessions across three years. Estimated glucosinolate content ranged from 11.0 to 35.8 micromoles per gram of fresh weight with an average value of 16.3 across three years. After that, a difference in glucosinolate content between winter and spring ecotypes was evaluated by ANOVA for three years as well as for mean values (Figure 4.3.1).

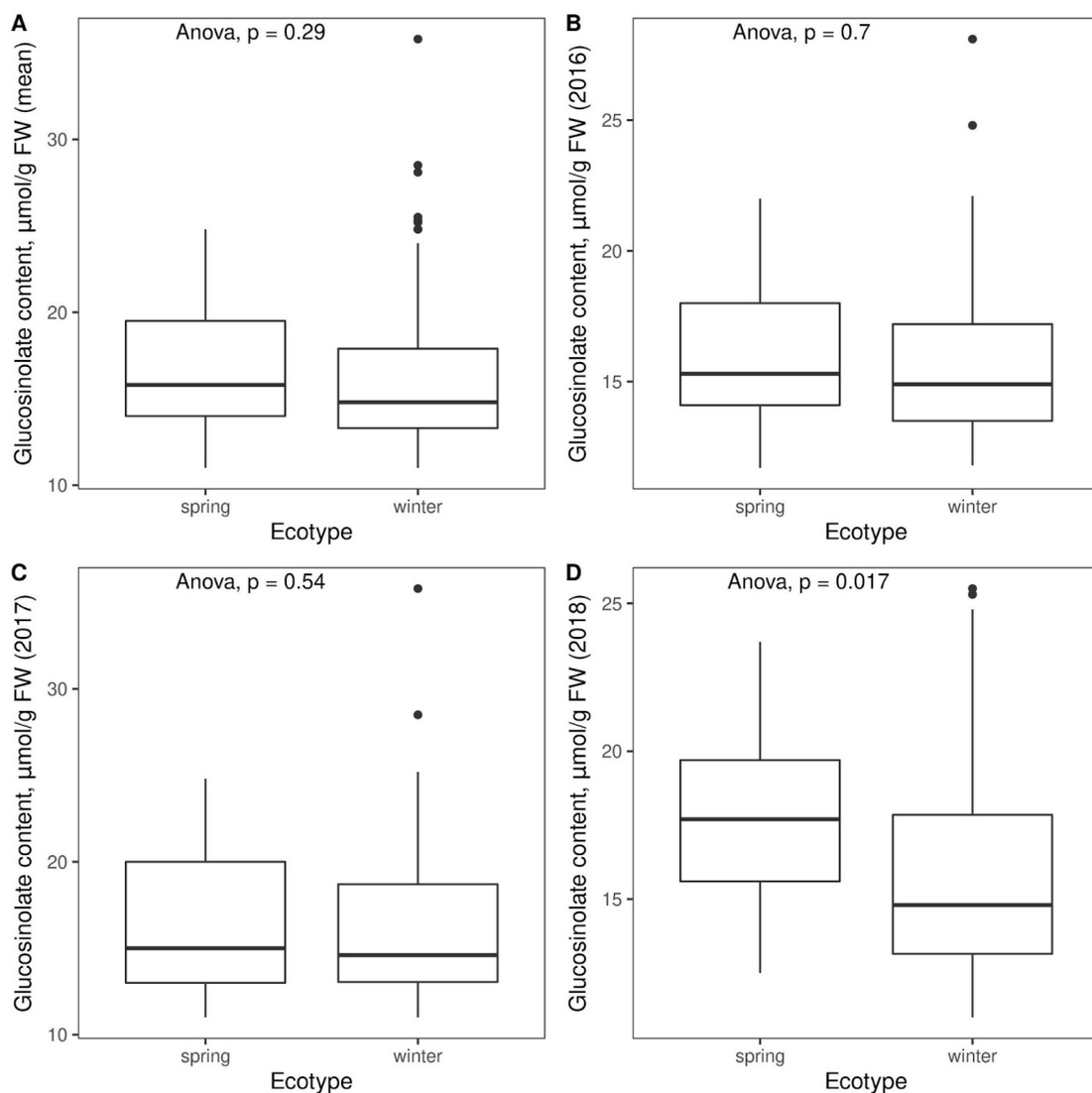


Figure 4.3.1 – Boxplots reflecting the distribution of the glucosinolate content across spring and winter accessions through 2016, 2017 and 2018.

This is due to the fact that it was previously demonstrated that quantitative traits such as yield, oil content, and height differences between the different rapeseed ecotypes (Assefa et al., 2018; Fridrihsone et al., 2020). It was demonstrated that no significant difference between the winter and spring phenotypes was identified at a 5% significance level for observations made in 2016 and 2017. Slight differences were only observed in 2018. Next, we tested the concordance of glucosinolate measurements across three years (Figure 4.3.2).

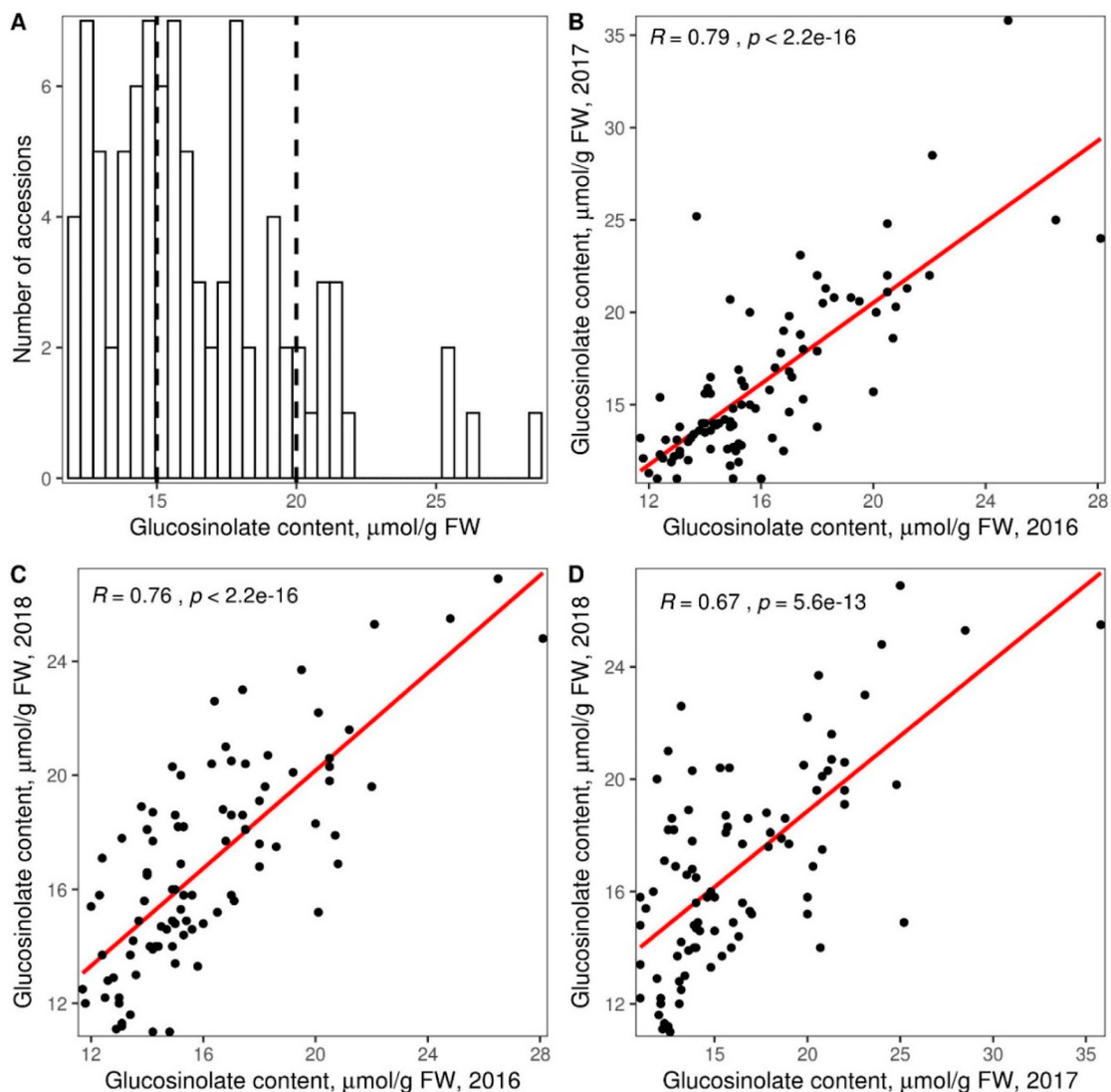


Figure 4.3.2 – Glucosinolate content distribution in the studied rapeseed cohort. Histogram depicting average glucosinolate content values for three years (A). Dashed lines divide low, middle, and high glucosinolate lines according to the (VNIIMK)

classification. Correlation of glucosinolate content for three vegetational seasons (B–D). Each dot corresponds to a plant sample. Regression lines are shown in red.

The glucosinolate content correlated positively and significantly between years, with Pearson’s correlation coefficient ranging from 0.67 to 0.79 (Pearson correlation, $p < 0.001$; Figure 4.3.2 A–C).

4.4 Association mapping and scanning for novel candidate genes

After the genotypes and phenotypes were collected, a scanning for genetic loci responsible for glucosinolate content was performed by using an MLM with additional PCs and kinship for fixed and random effects, respectively, to account for confounding derived from population structure and kin relationship. Since there were three years of observation, we applied an MLM for each year independently, after that a cumulative significance of genetic associations was calculated using Irwin-Hall distribution. After the application of the Bonferroni correction, two SNPs located 7 (A7) remained above the threshold (Figure 4.4.1, Figure 4.4.2).

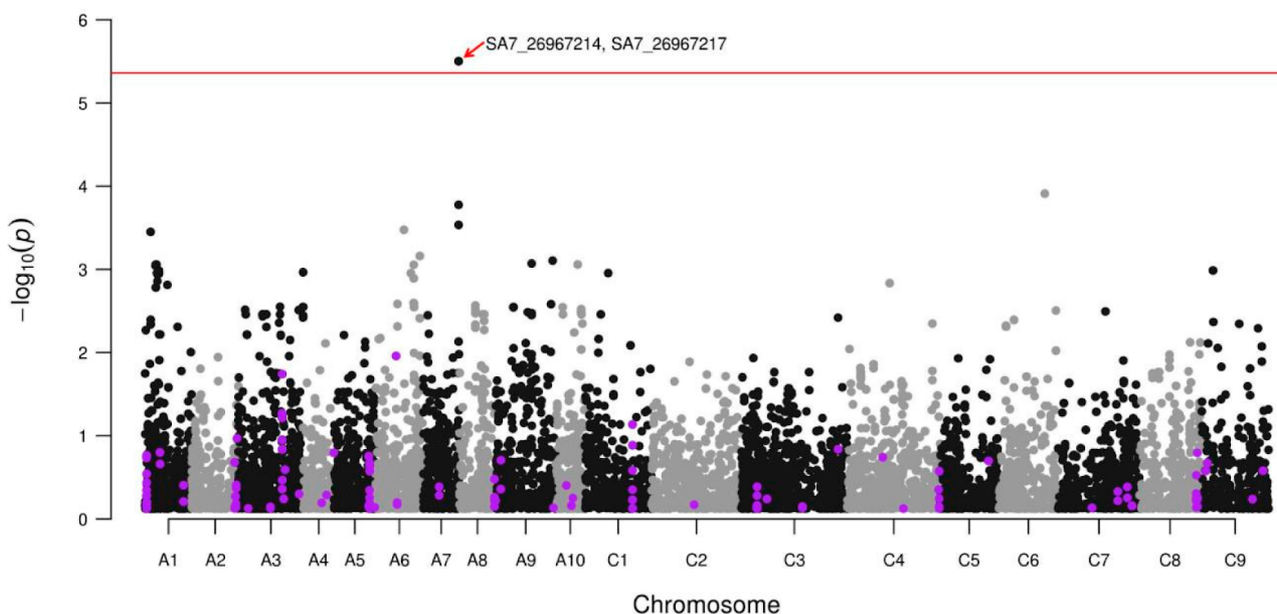


Figure 4.4.1 – Manhattan plot showing SNP markers associated with glucosinolate content. Each dot corresponds to a single SNP. Red line corresponds to the Bonferroni adjusted significance threshold. Purple dots correspond to SNPs

demonstrated to be significantly associated with glucosinolate content in the previous studies.

These two SNPs (SA7_26967214 and SA7_26967217) explained from 13.8 to 20.4 % of phenotype variance across three-year observations (Table 4.4.1). Notably, both demonstrated stability across three years (ANOVA p-value for SNP:Year = 0.899).

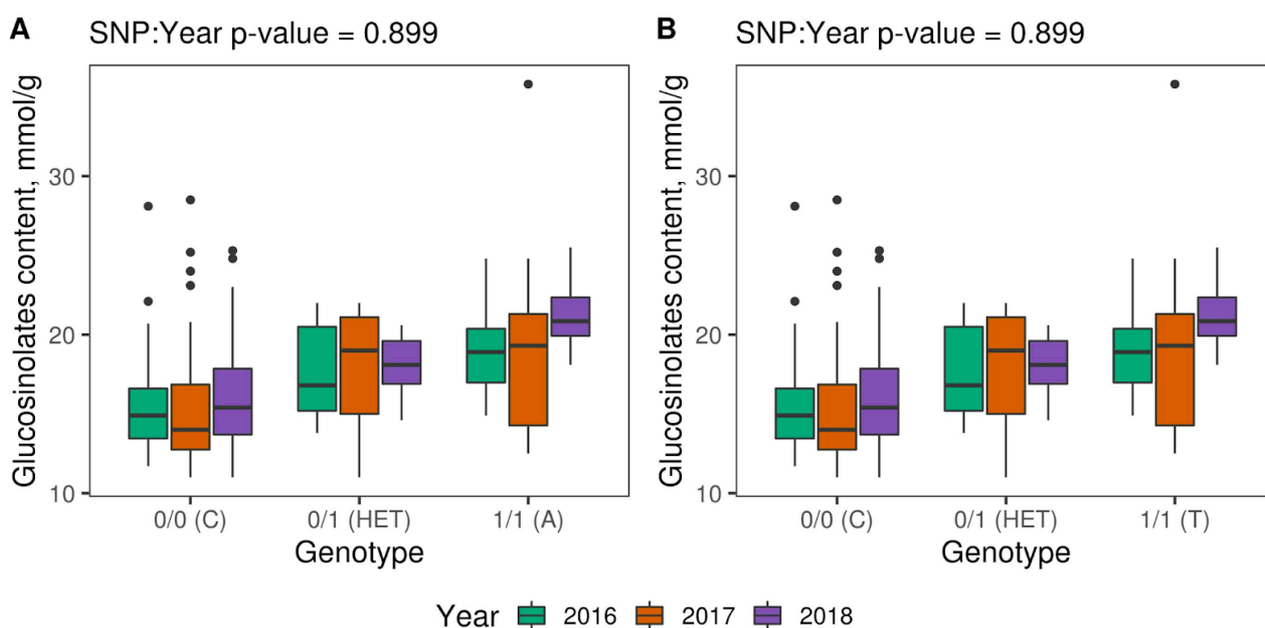


Figure 4.4.2 – Boxplots demonstrating the stability of the two significant SNPs across three years. Panel A corresponds to SA7_26967214, and panel B corresponds to SA7_26967217. Genotypes identified with 0/0, 0/1, 1/1 correspond to reference, heterozygous and alternative genotypes respectively, with corresponding nucleotides indicated in brackets. Colors correspond to the years.

A softer threshold of 0.0005 was also applied to find the additional genetic markers, as a result, additional five SNPs were detected that explained from 8.5 to 23.4% of glucosinolate content variation. In the previous studies it was demonstrated that the identified genetic markers explained 4 to 42% of phenotypic variance (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020; C.-M. Qu et al., 2015), this indicates

that the effects identified in the present study are in the concordance with the previously detected range.

Table 4.4.1 – List of SNPs significantly associated with the glucosinolate content.

SNP	PVE (2016)	P-value (2016)	PVE (2017)	P-value (2017)	PVE (2018)	P-value (2018)	P-value (Irwin-Hall)
SA1_4407039	0.113	0.0136	0.117	0.0127	0.234	0.0001	0.0003
SA6_21541176	0.127	0.0079	0.112	0.0176	0.202	0.0002	0.0003
SA7_26967211	0.092	0.0168	0.181	0.0003	0.141	0.0011	0.0001
SA7_26967214	0.204	0.0002	0.138	0.002	0.172	0.0002	3.15E-06
SA7_26967217	0.204	0.0002	0.138	0.002	0.172	0.0002	3.15E-06
SA7_26967241	0.085	0.0229	0.191	0.0002	0.143	0.0009	0.0002
SC6_35450938	0.107	0.0035	0.083	0.0083	0.089	0.0038	0.0001

Since it was demonstrated that the genetic diversity of the VNIIMK lines significantly differs (Chapter 4.2), the comparison between the previously identified SNPs (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020; C.-M. Qu et al., 2015; D. Wei et al., 2019) were compared to SNPs from the present study. To do so, SNPs associated with glucosinolate content for which the physical location was known were located in the genomic assembly used in the present study within the frame of 100kb. As a result, no significant associations for SNPs from the previous studies were identified. This could be explained by the difference in genetic background identified by PCA. Additionally, a smaller amount of the significant SNPs identified may be due to the fact that previous studies included larger amounts of accessions demonstrating broader trait variance of glucosinolate content ranging from 8 to 146 micromoles per gram of fresh weight (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020).

After the genetic associations were identified, scanning for potential candidate genes was performed within the 100kb region upstream and downstream of the

significant SNPs. Such distance was selected as the r^2 for this region remained high according to the LD analysis performed for the studied cohort r^2 value equalling 0.25 or more was specific for more than 30% of SNP pairs located within the 100 kb frame (Figure 4.4.1). Additionally, the same distance was used in the previous LD mapping studies in rapeseed to identify candidate genes (Guan et al., 2019; Q. Zhou et al., 2018; Q. Zhu et al., 2019).

Two significant SNPs SA7_26967214 and SA7_26967217 were localized within the intergenic region of genes for the U-box domain-containing protein 35-like and MTERF2 chloroplast-like. Both genes were not previously discussed in terms of regulation of glucosinolate concentration. Further analysis of the adjacent 100kb region revealed gene encoding BES1/BZR1 homolog protein 4-like located 75.3 kb downstream of the significant SNPs. This BES1 of the BZR family proteins was previously shown to be involved in brassinosteroid-dependent signaling and regulating the glucosinolate biosynthesis pathway (Guo et al., 2013). Additionally, 38.1 kb downstream of the significant SNPs, a gene encoding histone acetyltransferase HAC1 was identified. Genes encoding HAC1 histone acetyltransferase regulate senescence of the leaves, additionally it was demonstrated that the knockout mutants of *Arabidopsis thaliana* show down-regulation of genes encoding proteins involved in glucosinolate biosynthesis (Hinckley et al., 2019).

Of two other significant SNPs at $p = 0.0005$ the first one (SA6_21541176) represented a synonymous mutation within the gene encoding the derlin-2.1 protein. The second one (SA6_21541176) was annotated as a missense mutation within the gene of uncharacterized protein BNAA01G06520D. The other two SNPs are localized outside the gene regions. The last one (SA1_4407039) was located 31.8 kb downstream of the gene encoding the γ -glutamyl peptidase 1-like protein. It was previously shown that γ -glutamyl peptidases are involved in the biosynthesis of glucosinolates. Furthermore, *A. thaliana* plants with impaired genes encoding γ -glutamyl peptidases GGP1 and GGP3 demonstrated stained glucosinolate biosynthesis compared to the wild-type plants (Geu-Flores et al., 2009, 2011).

4.5 Conclusions

In the present part of the work a genetic collection of rapeseed accessions from the VNIIMK collection has been characterized. To do so we applied a GBS which allowed us to collect information on polymorphisms for genetic diversity description and association mapping at a reasonable time and financial costs.

Both ADMIXTURE and PCA identified significant population structure in the studied cohort. A clear separation previously identified between the spring and winter ecotypes (Gazave et al., 2016; D. Wei et al., 2017; D. Wu et al., 2019) was confirmed for the VNIIMK rapeseed collection. Additionally, a strong difference was revealed between yellow and dark seeded winter rapeseeds that was not previously reported and in the present study could be explained by the divergence in recent breeding history. Linkage disequilibrium analysis supported the previous observation of the higher LD in the C subgenome compared to the A subgenome (Qian et al., 2014; D. Wei et al., 2017; M. Zheng et al., 2017) which is explained by the larger size of C subgenome chromosomes compared to ones from A subgenome (F. Sun et al., 2017). These observations suggest that the rapeseed peculiarities of population structure are common for international and VNIIMK lines.

The scanning for genetic loci revealed two genetic markers significantly associated with the glucosinolate as well as an additional five marginally significant SNPs. The proportion of the variance explained by these markers was compared to the previously studied (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020; C.-M. Qu et al., 2015). Thus these genetic markers could be involved in the development of marker-assisted breeding in VNIIMK.

A scanning for candidate genes allowed to identify novel genetic loci associated with glucosinolate content. Notably, these ones were not previously identified by GWAS and QTL-based studies, which could be the result of studying the collections that differed from the VNIIMK collection at the genetic and phenotypic level. Previously a genetic markers associated with glucosinolate content was identified close to genes encoding enzymes involved in the biosynthesis of

glucosinolates. Specifically, MAM1 and MAM3 enzymes (Jan et al., 2019; Kittipol et al., 2019) are involved in glucosinolate side chain modification in *Arabidopsis* as well as AOP3 acting at the final stages of aliphatic glucosinolates biosynthesis (D. Wei et al., 2019). Additionally, genetic markers are located in the genes homologous to ones encoding enzymes for glucosinolate transporters GTR2 (S. Liu et al., 2020; C.-M. Qu et al., 2015). Several genetic markers strongly associated with glucosinolate content were located close to the genes encoding transcription factors of MYB28 and MYB34 families and transcription factor HAG1 controlling aliphatic glucosinolate biosynthesis (F. Li et al., 2014; D. Wei et al., 2019). In the present study SNPs marking the regions that were previously associated with the glucosinolates were not detected. This could be the result of the genetic difference that was identified while comparing VNIIMK and international lines. Additionally, the phenotypic diversity (glucosinolate concentration) was lower compared to the cohorts previously used to perform GWAS for that trait (Jan et al., 2019; F. Li et al., 2014; S. Liu et al., 2020).

Notably, several loci that were identified in the present study by means of GWAS were previously reported to be involved in the control of glucosinolates. Namely, by GWAS we additionally indirectly supported the role of γ -glutamyl peptidases, HAC1 histone acetyltransferase, BES1 of the BZR family proteins that were previously demonstrated to be involved in the regulation of the glucosinolate content control (Geu-Flores et al., 2009, 2011; Guo et al., 2013; Hinckley et al., 2019). Additionally, we identified novel genetic regions, containing genes that were not previously described, related to the regulation of glucosinolate content regulation which could facilitate a better understanding of the genetic control of glucosinolate content in rapeseed.

Chapter 5. QTL-mapping of oil-quality traits in sunflower

5.1 Introduction

Two traits associated with the oil quality in sunflower include tocopherol composition and oleic acid. Tocopherols are natural antioxidants that serve as a source of vitamin E. There are four types of tocopherols: α , β , γ , and δ . Notably, antioxidant activity increases in the row from α to δ while vitamin E activity decreases in the row (J. M. Fernández-Martínez et al., 2007). It should be mentioned that the natural diversity of tocopherols in sunflower is low and normally sunflower produces only α form (Y. Demurin et al., 1996; Hass et al., 2006). Thus several attempts at screening large genetic collections have been made to identify the mutations leading to the altered tocopherol composition. Such mutations were identified in sunflower lines from the Vavilov Institute (VIR) in Russia (Y. Demurin, 1993) and the Institute of Sustainable Agriculture (CSIC) collection in Spain (Velasco, Domínguez, et al., 2004). The mutations identified in the VIR collections were transferred to the LG15 and LG17 lines that are now part of VNIIMK genetic collection (Y. Demurin, 1993; Y. Demurin et al., 1996; Y. N. Demurin et al., 2004).

It was demonstrated that the tocopherol composition is controlled by two major effect loci *Tph1* (*m*) and *Tph2* (*g*), which are independently inherited (Y. N. Demurin et al., 2004). Using the CSIC lines and QTL mapping approaches *Tph1* was mapped on linkage group (chromosome) 1 and *Tph2* was mapped on linkage group 8 (Velasco, Pérez-Vich, et al., 2004; Vera-Ruiz et al., 2005). Mutation in *Tph1* was shown to be associated with the gene encoding 2-methyl-6-phytyl-1,4-benzoquinone (MPBQ) methyltransferase (MPBQ-MT) which converts MPBQ to 3,2-dimethyl-6-phytyl-1,4-benzoquinone (DMPBQ) (Tang, Hass, et al., 2006). The accumulation of the DMPBQ or MPBQ leads to the accumulation of the γ - and δ -tocopherols, respectively (Figure 5.1.1).

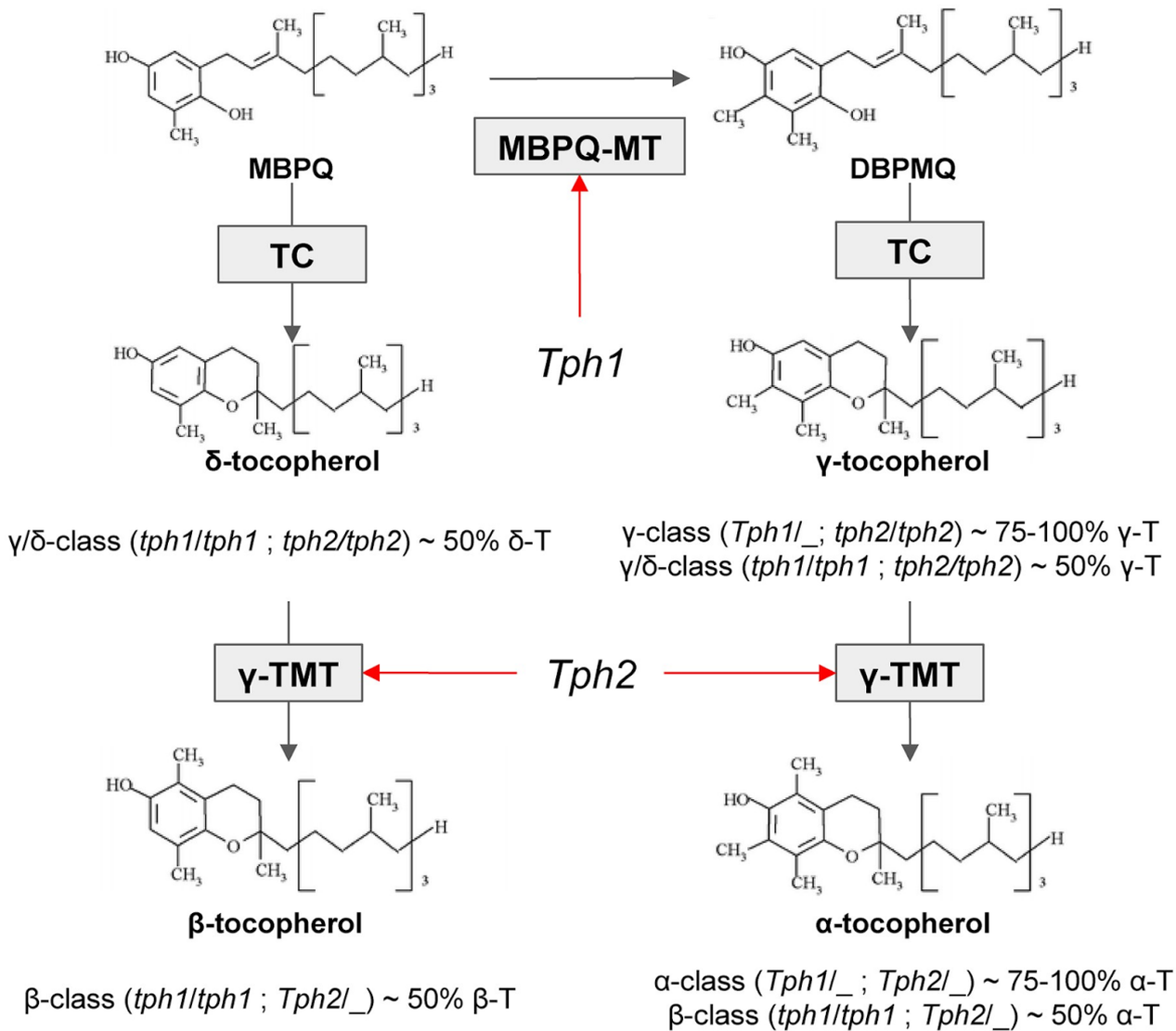


Figure 5.1.1 – Tocopherol biosynthesis scheme with corresponding phenotype classes. Names of the metabolites are indicated in bold: 2-methyl-6-phytyl-1,4-benzoquinone—MPBQ; 3,2-dimethyl-6-phytyl-1,4-benzoquinone — DMPBQ. Gray arrows correspond to the reactions catalyzed by the enzymes indicated in the gray squares: MPBQ methyltransferase—MPBQ-MT; tocopherol cyclase—TC; γ -tocopherol methyltransferase— γ -TMT. Red arrows indicate enzymes encoded by *Tph1* and *Tph2*. The biosynthesis pathway scheme was adapted from the previously published review (Lushchak & Semchuk, 2012).

In turn, the impaired *tph1* locus was associated with the increased accumulation of β -tocopherol. Mutation in *Tph2* was associated with the gene

encoding γ -tocopherol methyltransferase (γ -TMT), which converts γ - and δ -tocopherols to α - and β -tocopherols (Figure 5.1.1), respectively (Hass et al., 2006). Mutant *tph2* loci were associated with the accumulation of the γ -tocopherols. It also should be noted that the double mutant (*tph1/tph1; tph2/tph2*) plants accumulate γ - and δ -tocopherols in an almost equal ratio (Demurin, Borisenko, et al., 2006; Demurin et al., 2016; Gubaev et al., 2022).

Although it is assumed that the tocopherol composition is controlled by these two major effect loci, several studies identified additional and minor effect loci. Namely, a locus named *d* carrying gene coding for MPBQ-MT transferase (MT-2) paralogous to *m* (*Tph1*) was shown to interact epistatically with *m* (*Tph1*) and *g* (*Tph2*). This locus was mapped on chromosome 4 (Hass et al., 2006). Minor effect loci (modifying genes) affecting *tph2* mutation were mapped to chromosomes 1, 9, 14 and 16 (García-Moreno et al., 2012a). It was also assumed that the expressivity of *tph1* and *tph2* may depend on the genetic background of the lines (Y. Demurin et al., 1996). Thus the control of tocopherol composition in sunflowers remains under discussion. Additionally, currently, no genetic markers were identified for modern lines from the VNIIMK collection carrying *Tph1* and *Tph2* mutations.

Oleic acid is the monounsaturated fatty acid that on the one hand protects the lipids against thermooxidation during frying and on the other hand has positive health effects, namely lowering cholesterol and reducing inflammation (Sales-Campos et al., 2013). It should be noted that the antioxidant properties of oleic acid are increased in the oils with high content of γ - and/or δ -tocopherols (Skorić et al., 2008; Warner et al., 2008). Thus many breeding efforts in oilseeds including sunflower are made with the aim of increasing oleic acid content, especially in varieties used for frying oil production (Dehmer & Friedt, 1998; Monteros et al., 2008; Schierholt et al., 2000). As with tocopherols, there is low natural diversity for oleic acid in sunflowers (Cvejić et al., 2014). The first sunflower variety with high oleic acid content called Pervenets was obtained by chemical mutagenesis (Soldatov, 1976). Lately, this mutation was introduced within various collections with the aim of obtaining high-oleic varieties. It

was first assumed that the high oleic trait is controlled by a single dominant *Ol* gene (Urie, 1985). Studies based on hybridological analysis suggested more complex trait inheritance, in particular, it was assumed that this gene could be controlled by three dominant complementary genes (*O11-O13*) and one modifier gene (J. Fernández-Martínez et al., 1989; J. F. Miller et al., 1987). Development of the QTL mapping techniques allowed the identification of RAPD markers associated with the major *Ol* gene (Dehmer & Friedt, 1998). Lately, this gene explaining up to 56% of phenotype variance was mapped to chromosome 14 (Pérez-Vich et al., 2002). It was demonstrated that *Ol* is associated with the FAD2-1 gene encoding fatty acid desaturase that converts oleic acid to linoleic acid (Schuppert et al., 2006). A more recent study confirmed the role of *Ol* and identified additional minor effect loci located on chromosomes 8 and 9, explaining up to 10% of phenotypic variance (Premnath et al., 2016). A QTL study based on high-throughput genotyping additionally identified loci located in linkage groups 9 and 6, explaining 12 and 6% of phenotype variance (F. Zhou et al., 2018). These observations are in concordance with the hypothesis on the effect of genetic background on the *Ol* gene that has been previously put forward (Demurin, Skoric, 1996). Thus as in the case of tocopherols, the genetic control of oleic acid remains under discussion and similarly, there are no genetic markers that have been found with the implementation of lines from Russian genetic collections.

Thus the aim of this part of the research was to identify loci and perspective SNP markers associated with altered tocopherol composition and high oleic acid content. To do so we analyzed two experimental crosses derived from crossing contrast modern lines from VNIIMK. The part of the results related to the genetic map construction and mapping of the tocopherol composition presented in the following chapter was published in the G3-Genes Genomes Genetics journal (Gubaev et al., 2022).

5.2 Phenotype evaluation

To characterize tocopherol composition a thin-layer chromatography (TLC) was applied. As a result 142 and 144 F2 seeds for crosses VK195xVK303 and VK876xVK101 were phenotyped, respectively. Parental lines were phenotyped in at least 7 replicates. Maternal lines VK195 and VK876 were attributed to the γ/δ phenotype as they demonstrated almost equal γ -/ δ -tocopherol content. Paternal lines VK303 and VK101 demonstrated a wild-type α phenotype; the proportion of α -tocopherol equaled 100% (Table 5.2.1).

Table 5.2.1 – Tocopherol composition in parental lines.

Line name	Genotype	Tocopherol class	Mean proportion of α -tocopherol, %	Mean proportion of β -tocopherol, %	Mean proportion of γ -tocopherol, %	Mean proportion of δ -tocopherol, %
BK195	<i>tph1/tph1;tph2/tph2</i>	γ/δ	0	0	51.7 \pm 5.41	48.3 \pm 5.41
BK303	<i>Tph1/Tph1;Tph2/Tph2</i>	α	100	0	0	0
BK876	<i>tph1/tph1;tph2/tph2</i>	γ/δ	0	0	52.9 \pm 8.71	47.1 \pm 8.71
BK101	<i>Tph1/Tph1;Tph2/Tph2</i>	α	100	0	0	0

F2 progeny was classified into tocopherol phenotypic classes based on the tocopherol composition depending on the putative allelic states of two major genes *Tph1* and *Tph2*. Each of the plants was assigned to one of the four tocopherol classes. As a result, 142 and 144 phenotyped F2 seeds from the VK195xVK303 and VK876xVK101, 137 (96.4%) and 143 (99.3%) were classified into the known phenotypic classes, respectively (Figure 5.2.1).

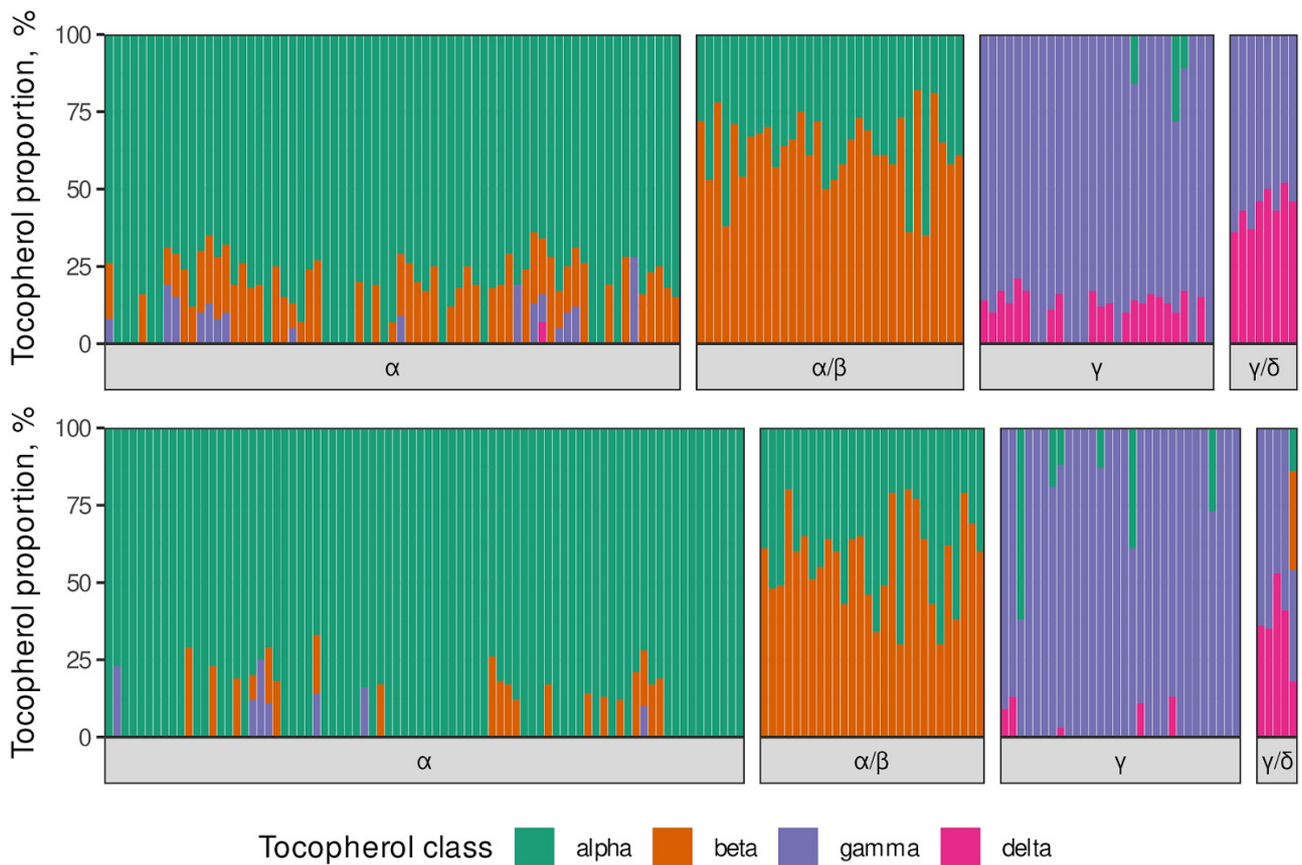


Figure 5.2.1 – Relative content of each of the tocopherol classes among the genotyped individuals in the F2 progeny. Each bar corresponds to a single F2 seed. Each colored bar shows the proportion of each of the four tocopherol classes. Facets show the distribution of phenotyped F2 seeds across the α -, α/β -, γ -, and γ/δ -tocopherol phenotypic classes. The upper panel corresponds to the VK195xVK303 cross. The lower panel corresponds to the VK876xVK101 cross.

Only one and five F2 seeds from VK195xVK303 and VK876xVK101 crosses, respectively, were not assigned to any specific phenotypic class since the tocopherol ratios for these accessions deviated from the specified classes. Phenotype classes of the classified F2 progeny matched the 9:3:3:1 distribution according to the χ^2 goodness-of-fit test (Table 5.2.2) with high confidence (P-values of 0.4505 and 0.5382 for VK195xVK303 and VK876xVK101, respectively). These observations speak in favor of a two-gene model.

Table 5.2.2 – Number of plants assigned to one of four tocopherol classes for each of the crosses.

Cross	α -class	α/β -class	γ -class	γ/δ -class	P-values for χ^2 goodness of fit test
VK195xVK303	69	32	28	8	0.4505
VK876xVK101	80	28	30	5	0.5382

To characterize relative oleic acid content, a gas chromatography followed by mass spectrometry was applied. As a result, the relative content of oleic acid was determined for parental lines in at least seven replicates and for 142 and 144 F2 seeds for crosses VK195xVK303 and VK876xVK101, respectively (Figure 5.2.2).

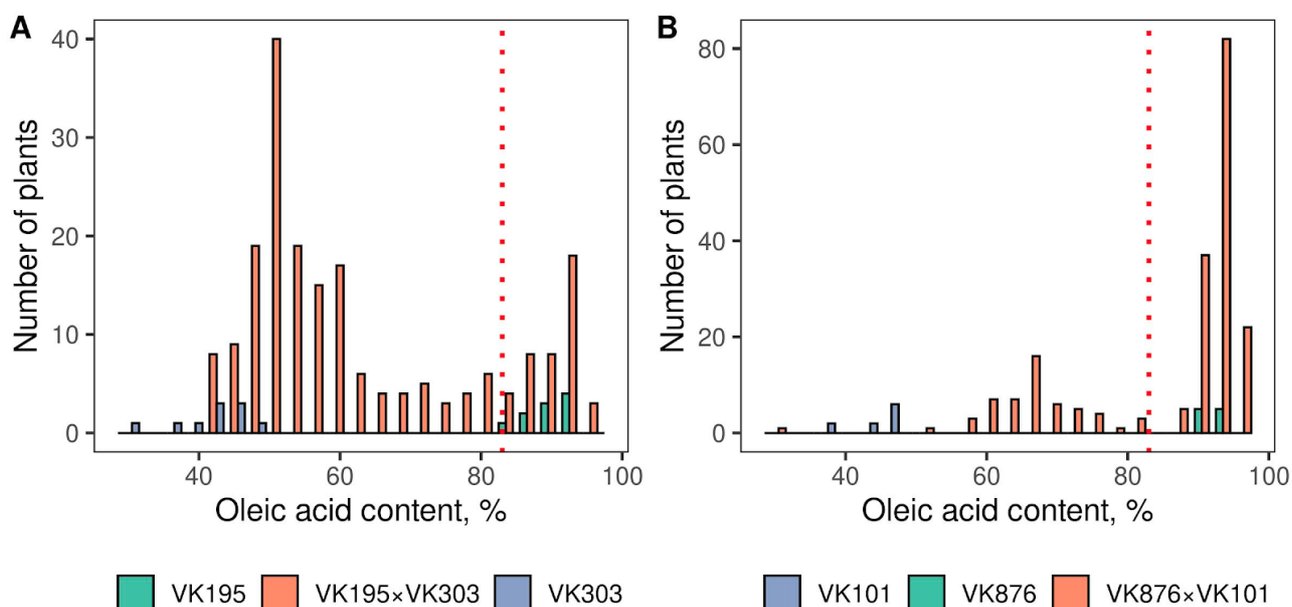


Figure 5.2.2 – The distribution of the oleic acid content for parental lines and F2 progeny. Panel A reflects the distribution of oleic acid content for cross VK195xVK303, panel B reflects the distribution of oleic acid content for cross

VK876xVK101. The green color of the bars corresponds to wild-type parental lines, the blue color corresponds to mutant parental lines, and the orange color corresponds to F2 progeny. Dotted red lines correspond to oleic acid content specific for high-oleic plants.

As the major effect of the dominant *Ol* gene was assumed for studied populations, plants were distributed to high-oleic and non-high-oleic classes (Table 5.2.3).

Table 5.2.3 – The distribution of the F2 progeny into high-oleic and non-high oleic classes.

Cross	High-oleic	Non-high-oleic	P-values for χ^2 goodness of fit test
VK195xVK303	40	104	0.1461
VK876xVK101	114	28	< 2.2e-16

For population VK195xVK303 a χ^2 goodness of fit test confirmed a 3:1 segregation ratio, where 3 parts correspond to the high-oleic class and 1 to non-high-oleic classes. For the population VK876xVK101 a χ^2 goodness of fit test didn't confirm the 3:1 segregation ratio (Table 5.2.3). In contrast, an opposite distribution was observed: 3 parts were represented by non-high-oleic and 1 part of high-oleic class (P-value of χ^2 goodness of fit test = 0.4414).

5.3 Genotyping and genetic map construction

For genetic map construction we applied a genotyping-by-sequencing approach to identify genetic polymorphisms for studied crosses. As a result, 425213 and 499063 raw SNPs were collected for VK195xVK303 and VK876xVK101, respectively. Next SNPs were filtered according to the filters (see methods section) as well as SNPs that were homozygous within and polymorphic between the parents were selected as this is a prerequisite for genetic map construction. As the low SNP coverage is a common problem for the GBS while being applied for F2 populations that demonstrate high amounts of heterozygous positions, an imputation based on the LB-Impute algorithm was applied (Fragoso et al., 2016). Imputation decreased the proportion of missing genotypes from 0.22 to 0.1 for the VK195xVK303 cross and from 0.2 to 0.09 for the VK876xVK101 cross. After imputation, 7028 SNPs for the VK195xVK303 cross and 5,876 SNPs for the VK876xVK101 cross were used for map construction (Figure 5.3.1) within the r/qtl package (Broman et al., 2003). During the genetic map construction genetic markers were additionally filtered by the expected genotype segregation pattern for F2 populations (1:2:1). Additionally, markers that significantly affected the map length were discarded (for details see methods section).

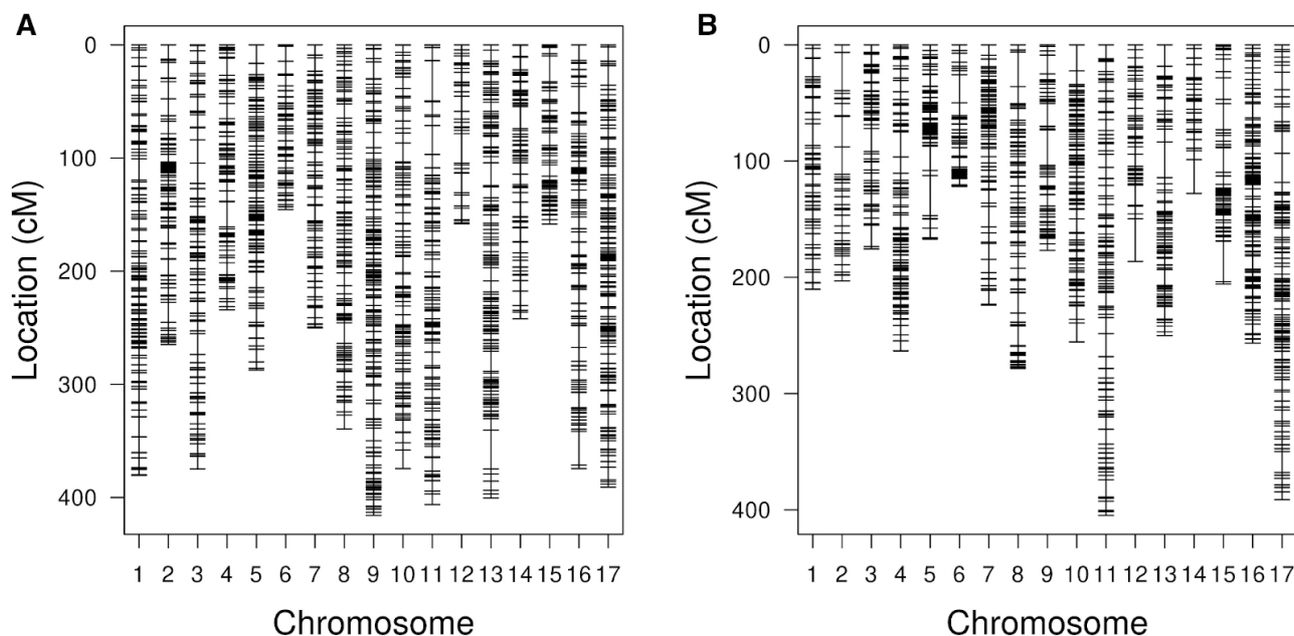


Figure 5.3.1 – SNP-based genetic linkage maps for the VK195xVK303 (panel A) and VK876xVK101 (panel B) crosses.

A genetic map for cross VK195xVK303 consisted of 3200 SNPs and spanned a total distance of 5,197.7 cM, with the maximum spacing of 35.5 cM and the average intermarker distance of 1.63 cM. The correlation between physical and genetic map was significant (P-value for Pearson correlation coefficient < 1.223024e-19), Pearson correlation coefficient ranged from 0.62 to 0.97 across the chromosomes (Table 5.3.1).

Table 5.3.1 – Summary for the genetic maps constructed for VK195xVK303 and VK876xVK101 crosses.

Chromosome	Cross	Number of markers	Map length (cM)	Average spacing (cM)	Maximum spacing (cM)	Pearson correlation coefficient	P-value For Pearson correlation coefficient
1	VK195 x VK303	189	380.54	2.02	18.87	0.9	6.01E-68
2		188	264.79	1.42	17.82	0.72	3.1E-31
3		163	374.72	2.31	20.43	0.86	7.8E-49
4		136	234.29	1.74	22.41	0.85	1.18E-39
5		246	287.46	1.17	16.34	0.82	3.18E-62
6		95	145.51	1.55	13.32	0.88	1.52E-32
7		173	250.03	1.45	13.48	0.62	1.22E-19
8		238	339.45	1.43	18.24	0.94	1.42E-109
9		315	415.84	1.32	14.74	0.97	1.15E-188
10		240	374.47	1.57	16.69	0.95	1.42E-119
11		194	406.14	2.1	35.49	0.62	2.14E-22
12		57	158.27	2.83	26.02	0.91	2.02E-22

13		216	400.4	1.86	34.37	0.93	4.96E-98
14		126	242.1	1.94	19.66	0.9	3.36E-46
15		127	158.39	1.26	22.72	0.91	1.79E-48
16		187	374.41	2.01	29.86	0.79	1.8E-41
17		310	390.89	1.27	17.23	0.92	1.2E-126
overall		3200	5197.71	1.63	35.49	-	-
1	VK876 x VK101	98	210.29	2.17	18.77	0.9	1.7E-36
2		77	203.15	2.67	32.93	0.93	7.75E-34
3		118	175.66	1.5	19.68	0.9	8.98E-45
4		190	263.26	1.39	21.52	0.96	3.44E-106
5		138	167.13	1.22	34.49	0.89	2.48E-49
6		104	121.91	1.18	24.18	0.89	1.13E-36
7		215	223.92	1.05	17.5	0.48	5.21E-14
8		184	278.34	1.52	29.87	0.92	1.92E-75
9		104	176.82	1.72	30.57	0.95	5.43E-54
10		225	255.63	1.14	22.49	0.89	1.58E-79
11		137	404.86	2.98	19.73	0.9	1.25E-49
12		100	186.25	1.88	36.6	0.81	3.54E-24
13		165	250.15	1.53	30.8	0.96	9.5E-90
14		75	127.86	1.73	29.11	0.9	1.52E-27
15		105	205.8	1.98	34.89	0.94	7.57E-50
16		236	256.59	1.09	12.07	0.87	2.24E-73
17		300	391.19	1.31	24.98	0.89	2.8E-102
overall		2571	3898.81	1.53	36.6	-	-

For cross VK876xVK101 genetic map consisted of 2571 SNPs and spanned 3,898.8 cM. With the maximum spacing of 36.6 cM and average intermarker distance of 1.53 cM. The correlation between physical and genetic maps was significant (P-value for Pearson correlation coefficient $< 5.205962e-14$) and Pearson correlation coefficient ranged from 0.48 to 0.96 across the chromosomes (Table 5.3.1).

Previously the high-density SNP based genetic maps based on the reduced representation sequencing (SLAF-, RAD-, GBS-sequencing) data for sunflower included from 817 to 6,136 SNPs and was from 1444 to 2472 cM in length (Celik et al., 2016; Talukder et al., 2014; F. Zhou et al., 2018). There are two possible explanations that in the present study genetic maps were almost two times larger compared to the previously reported ones. First, relatively relaxed filters were applied for segregation distortion. Second, errors in genotyping were present due to the nature of the sunflower genome which contains a high amount of repetitive regions (Badouin et al., 2017). This in turn may lead to errors during read alignment (Kane et al., 2011; Treangen & Salzberg, 2012). It also should be noted that for other plant species large genetic maps and large chromosomes compared to the obtained ones were also reported. Large chromosomes that were longer than 300 cM were reported for maize (Su et al., 2017; J. Wang et al., 2018) and wheat (Yang et al., 2018); both maps were based on the reduced representation sequencing. A large RAD-seq based genetic map was also reported for sweetpotato (Shirasawa et al., 2017); the map was 7313.5 cM in length with the largest chromosome of 904.5 cM. The correlation between the genetic and physical distance reported in the present study was higher compared to the previously reported values of the Pearson correlation coefficient ranging from 0.2 to 0.7 (Celik et al., 2016). This speaks in favor that the constructed genetic maps are in concordance with the physical ones.

5.4 QTL-mapping of tocopherol composition

The next stage of the study was related to the association mapping of tocopherol composition. Here two approaches have been implemented; the first one was associated with mapping the proportion of each tocopherol class as independent observation (quantitative mapping). The second was related to the mapping of the tocopherol composition associated with the putative allelic states of *Tph1* and *Tph2* revealed by the belonging of the plant phenotypes to one of the four tocopherol classes (qualitative mapping).

The quantitative mapping approach revealed loci associated with α - β -, γ - and δ -tocopherols for both crosses (Figure 5.4.1). Additionally, a proportion of phenotypic variance explained by SNP tagged loci was calculated for markers 1.5-LOD interval estimated for the most significant marker (Supplementary Table 4 from

<https://academic.oup.com/g3journal/article/12/4/jkac036/6527637#supplementary-data>). For cross VK195xVK303 loci associated with the proportion of α -tocopherol were located on chromosome 1, additionally, loci associated with a proportion α -, β -, γ - and δ -tocopherols were mapped on chromosome 8. The maximum proportion of variance explained by markers located on chromosome 1 explained 21.98% and 38.14% of α - and β -tocopherol variance. For cross VK876xVK101 loci associated with β -tocopherol were mapped on chromosome 1, additionally, loci associated with α -, γ -, and δ -tocopherols were mapped on chromosome 8. The maximum proportion of β -tocopherol variance explained by markers located on chromosome 1 was 44.44%. The markers located on chromosome 8 explained up to 71.09%, 85.74%, and 9.83% of α -, γ -, and δ -tocopherol content.

For both crosses associations for β -tocopherol were identified on chromosome 1 while associations for α -, γ -tocopherol were found on chromosome 8 which is in concordance with the previously published results (García-Moreno et al., 2012b; Tang, Hass, et al., 2006; Vera-Ruiz et al., 2006).

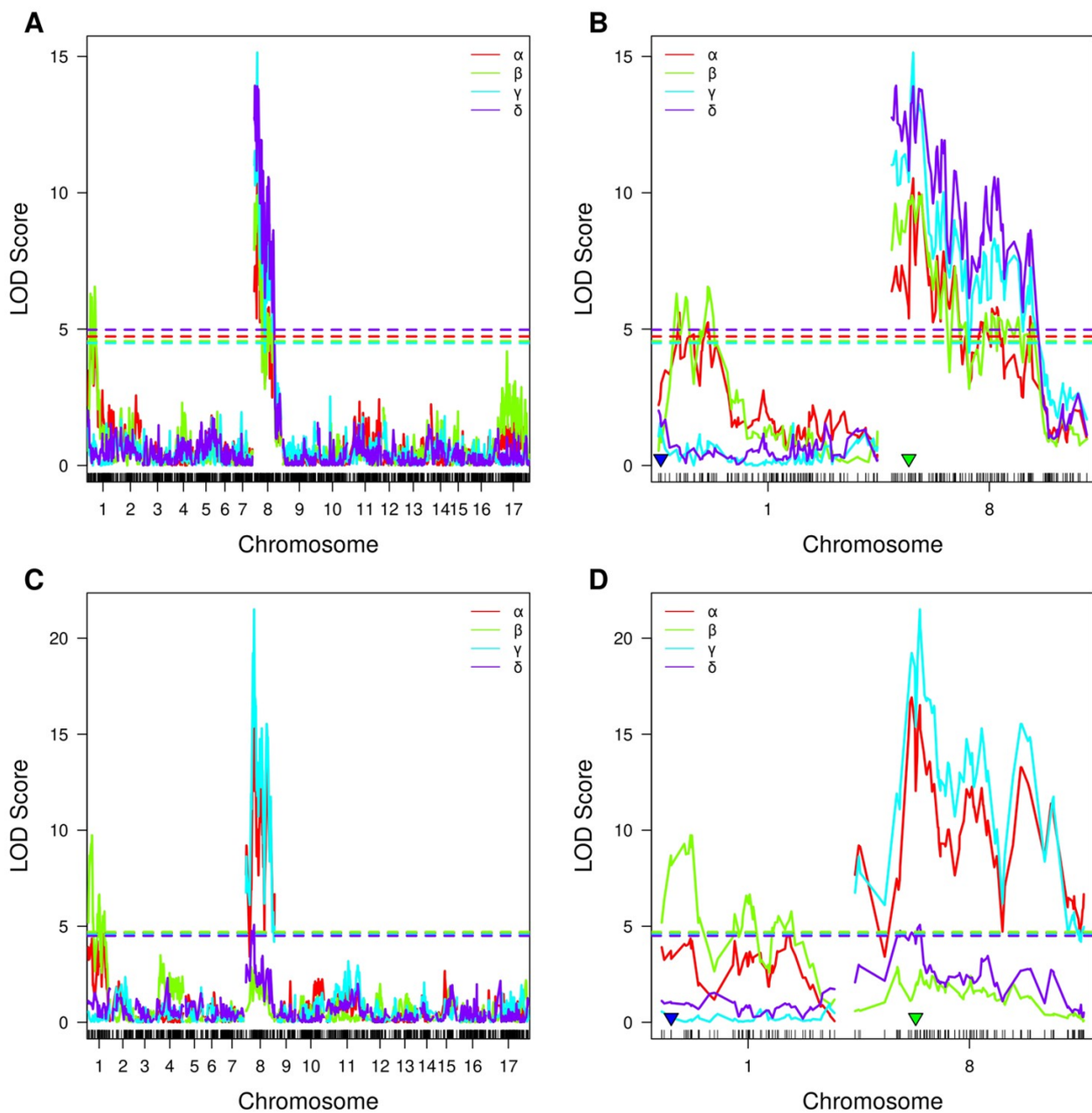


Figure 5.4.1 – Likelihood curve of the LOD score for α -, β -, γ -, and δ -tocopherol content for the VK195xVK303 a and b) and VK876xVK101 c and d) populations obtained by using the interval mapping approach. Dashed lines correspond to the permutation threshold. The color of lines corresponds to the tocopherol type. a and c) The mapping results for all chromosomes. b and d) The mapping results for chromosomes carrying significant markers. Blue and green triangles indicate markers that are most closely located to the *Tph1* and *Tph2* loci, respectively, based on the physical map data.

Additionally, the differences in mapping results were identified between the crosses which could be the result of different distributions of the relative content of tocopherol classes (Figure 5.2.1). Compared to the previous studies the proportion of variance explained by the markers was significantly higher, namely, for β - tocopherol genetic markers explained 90% of phenotypic variance (Vera-Ruiz et al., 2006) and for γ -tocopherols genetic markers explained 97% of the variance (García-Moreno et al., 2006). This could be explained by the fact that compared to the previous studies, double mutant lines were analyzed in which the tocopherol composition is controlled by two independent loci simultaneously. Furthermore, no additional loci that were previously mapped on chromosomes 4, 9, 14, 16 were not identified (García-Moreno et al., 2012b; Hass et al., 2006). This could be explained by the relatively small amount of progeny used to find associations as well as with the strong effect of major effect loci located on chromosomes 1 and 8 which mask minor effect loci. This fact makes double mutant lines VK195 and VK876 carrying *tph1* and *tph2* mutant alleles with high expressivity convenient for breeding programs (Y. N. Demurin et al., 2004). Thus the next step of the present study was directly related to the mapping of major effect *Tph1* and *Tph2* loci.

To do so we applied qualitative mapping of *Tph1* and *Tph2* as the distribution of the classes followed a 9:3:3:1 distribution. For each phenotype of the plant the potential presence or absence of *Tph1* and *Tph2* were assigned as phenotypes. The quantitative approach mapped *Tph1* within a 1.5-LOD interval spanning from 77.77 to 93.35 cM on chromosome 1 with the maximum LOD score of 20.19 for VK195xVK303 cross (Figure 5.4.2 A, B). For cross, VK876xVK101 *Tph1* was mapped on chromosome 1 from 3.05 to 44.54 cM 1.5-LOD interval with the maximum LOD score of 15.6 (Figure 5.4.2 C, D). For cross VK195xVK303 *Tph2* was mapped on chromosome 8 from 32.12 to 64.69 cM with a maximum LOD score of 18.33 (Figure 5.4.2 A, B), for cross VK876xVK101 *Tph2* associated region was located from 75.98 to 83.57cM on chromosome 8 with maximum LOD value equal to 24.65 (Figure 5.4.2 C, D).

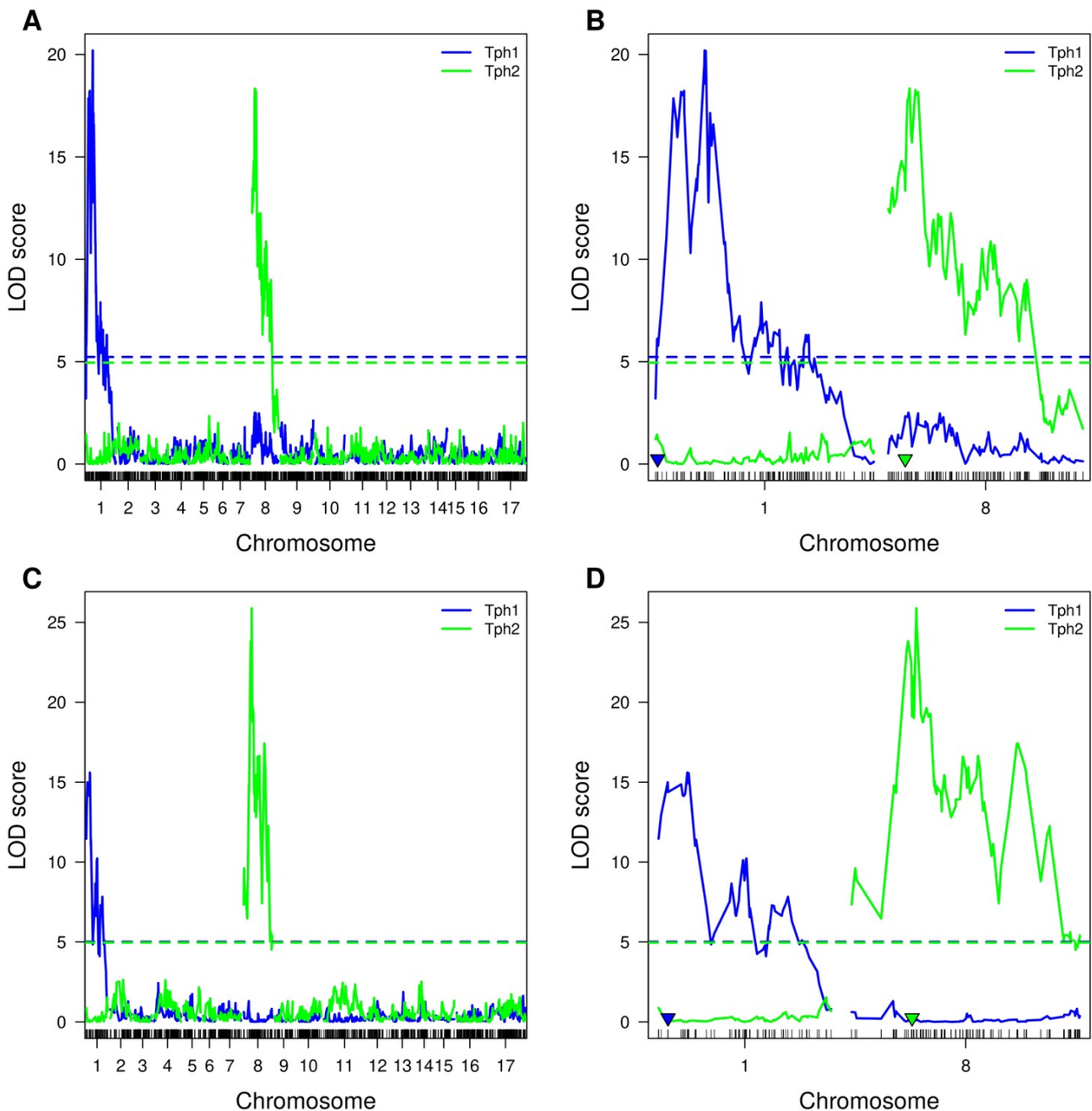


Figure 5.4.2 – Likelihood curve of the LOD score for *Tph1* and *Tph2* for the VK195xVK303 a and b) and VK876xVK101 c and d) populations. Dashed lines correspond to permutation results. Interval mapping results for *Tph1* and *Tph2* are indicated with the corresponding colors. a and c) The mapping results for all chromosomes. b and d) The mapping results for chromosomes carrying significant markers. Blue and green triangles indicate markers that are most closely located to the *Tph1* and *Tph2* loci, respectively, based on the physical map data.

Next, we scanned for the closest markers located to the causal genes associated with *Tph1* and *Tph2*. According to the information on the physical location of the *Tph1* gene encoding MPBQ-MT (NCBI Gene ID: 110937001) it was established that the closest significant markers were located 0.5 (marker S1_9228105) and 5.3 megabases (marker S1_4480460) for crosses VK195xVK303 and VK876xVK101, respectively. These markers demonstrated significant LOD scores of 6.11 and 14.37. For *Tph2* locus carrying genes encoding γ -TMT1 and γ -TMT2 (NCBI Gene IDs: 110872346 and 110872347) closest significant markers were located 0.34 (marker S8_21613153) and 0.42 megabases (marker S8_21536716), for crosses VK195xVK303 and VK876xVK101, respectively. The respective LOD scores were 13.34 and 19.10, respectively.

The results of mapping were in concordance with the published ones. Previously *Tph1* was also mapped on the upper end of chromosome 1 using F2 crosses (Tang, Hass, et al., 2006; Vera-Ruiz et al., 2006). The markers associated with *Tph1* were also physically located to the gene encoding MPBQ-MT. For *Tph2* it was also demonstrated that this locus is located on chromosome 8 for different mapping populations (García-Moreno et al., 2006; Hass et al., 2006). As in the case with *Tph1* markers, *Tph2*-associated ones were closely located to genes encoding γ -TMT1 and γ -TMT2 (Tang, Hass, et al., 2006). It should be noted that in the above-mentioned studies a quantitative approach was applied i.e. the mapping was performed for relative content of β - and γ -tocopherol proportions were analyzed (García-Moreno et al., 2006, 2006; Hass et al., 2006; Tang, Hass, et al., 2006). Notably, different statistical approaches were used to map these traits, including single marker ANOVA (García-Moreno et al. 2006; Hass et al. 2006; Vera-Ruiz et al. 2006) and interval mapping (García-Moreno et al. 2006), composite interval mapping (García-Moreno et al. 2012). Here, we have demonstrated that in our case, the qualitative approach based on interval mapping for traits with binary distribution was also suitable for mapping major effect loci and helped us to map *Tph1* and *Tph2* in both populations.

5.5 QTL mapping of oleic acid content

To map oleic acid content, three approaches have been used. First, the raw phenotypes were mapped, i.e the relative content of the oleic acid by two methods: interval mapping adapted for non-normally distributed traits and composite interval mapping as it was previously used to map this trait. Additionally, the transformation of the data into binary traits was performed following the mapping of high-oleic ($\geq 83\%$) and non-high oleic traits ($< 83\%$) as this threshold was previously used to divide high and non-high oleic phenotypes (Lacombe et al., 2009).

For cross VK195xVK303 a peak located on chromosome 14 was identified with all three methods (Figure 5.5.1). A 1.5-LOD confidence interval calculated for composite interval mapping results spanned from 199.76 to 208.4 cM. The markers that were located within the 1.5-LOD confidence interval explained up to 56.37% of phenotyping variance (Table 5.5.1, Figure 5.5.1). According to the physical map information, a genetic marker S14_137961667 was most closely (13.3 megabases away) located to the gene encoding FAD2-1 (NCBI GeneID 110904312) located from 151341021 to 151344179 bp and associated with high oleic acid content (Lacombe & Bervillé, 2001; Schuppert et al., 2006). Notably for the cross VK195xVK303 genetic markers for high content of oleic acid demonstrated the recessiveness, which was expected starting from the analysis of the phenotype distribution that was contrariwise compared to the expected 3:1 segregation ratio where 3 should correspond to high-oleic phenotypes and 1 part for non-high oleic phenotypes. The expected segregation was previously explained due to the dominant nature of the mutant *Ol* allele that carries two copies of gene encoding fatty acid desaturases, it was assumed that the one impaired copy FAD2-1 could reduce the expression of normal FAD2-1 via production of siRNA (Lacombe et al., 2009). The fact that the expected segregation ratio was not observed as well as the recessive nature of the genetic markers could be explained by the presence of the additional recessive *Ol* allele, and/or the effect of the genetic background of the wild-type line VK303 as the dominant state of the *Ol* was expected for the carrier line VK195.

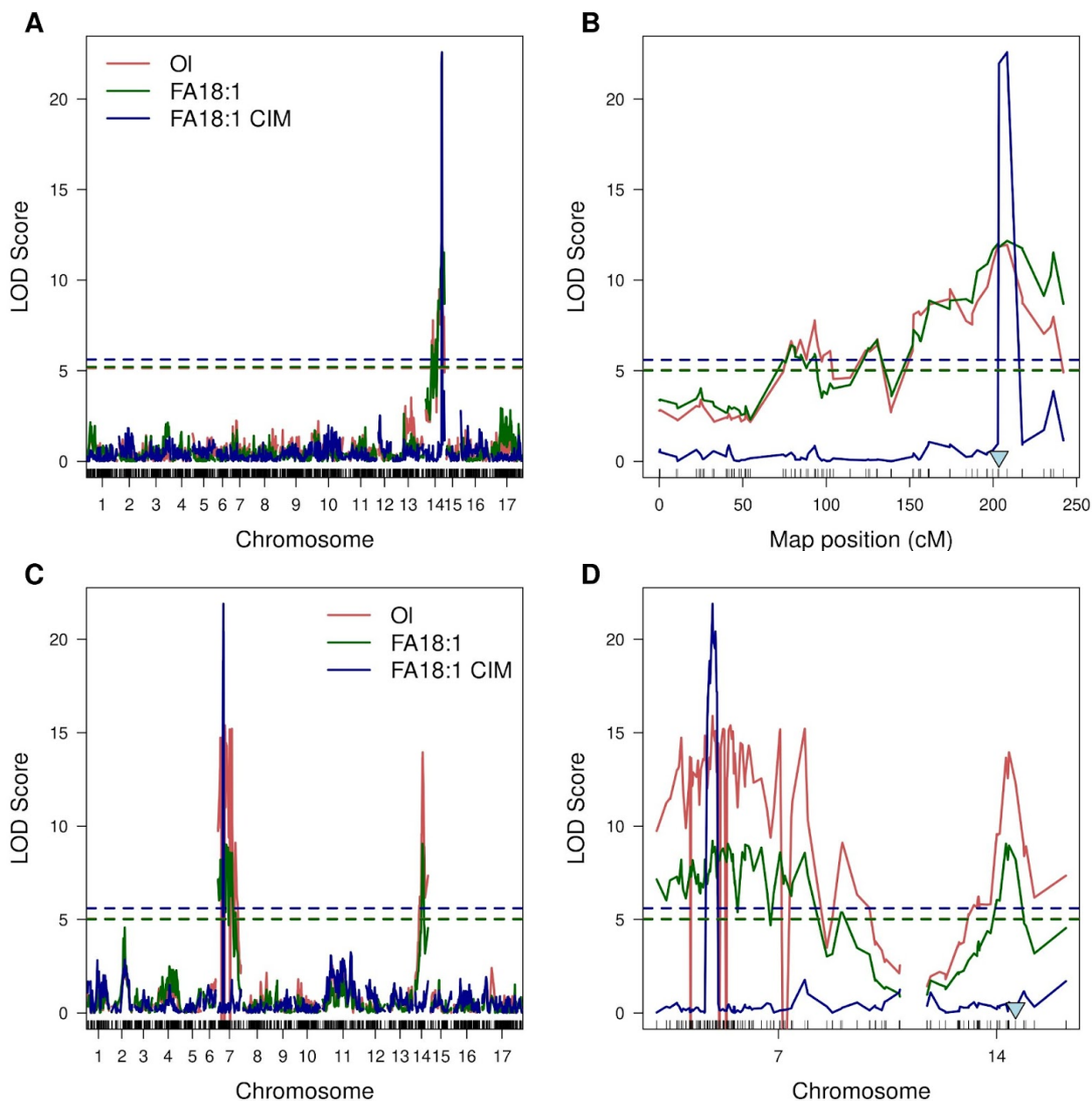


Figure 5.5.1 – Likelihood curves of the LOD score *Ol* gene and oleic acid content for VK195xVK303 (panels A and B) and VK876xVK101 (panels C and D) crosses. The red curve reflects the results of mapping the binary trait associated with the presence and/or absence of the *Ol* gene with the interval mapping approach. The green curve reflects the results of interval mapping adapted for non-normally distributed traits of oleic acid content. The blue curve reflects the results of composite interval mapping of oleic acid content. Light blue triangles indicate markers that are most closely located to the *Ol* gene encoding FAD2-1 based on the physical map data.

For cross VK876xVK101 interval mapping for both types of traits (binary and numeric) identified loci located on chromosomes 7 and 14 which contradicted the expectation of the presence of one major effect loci (Figure 5.5.1 C, D).

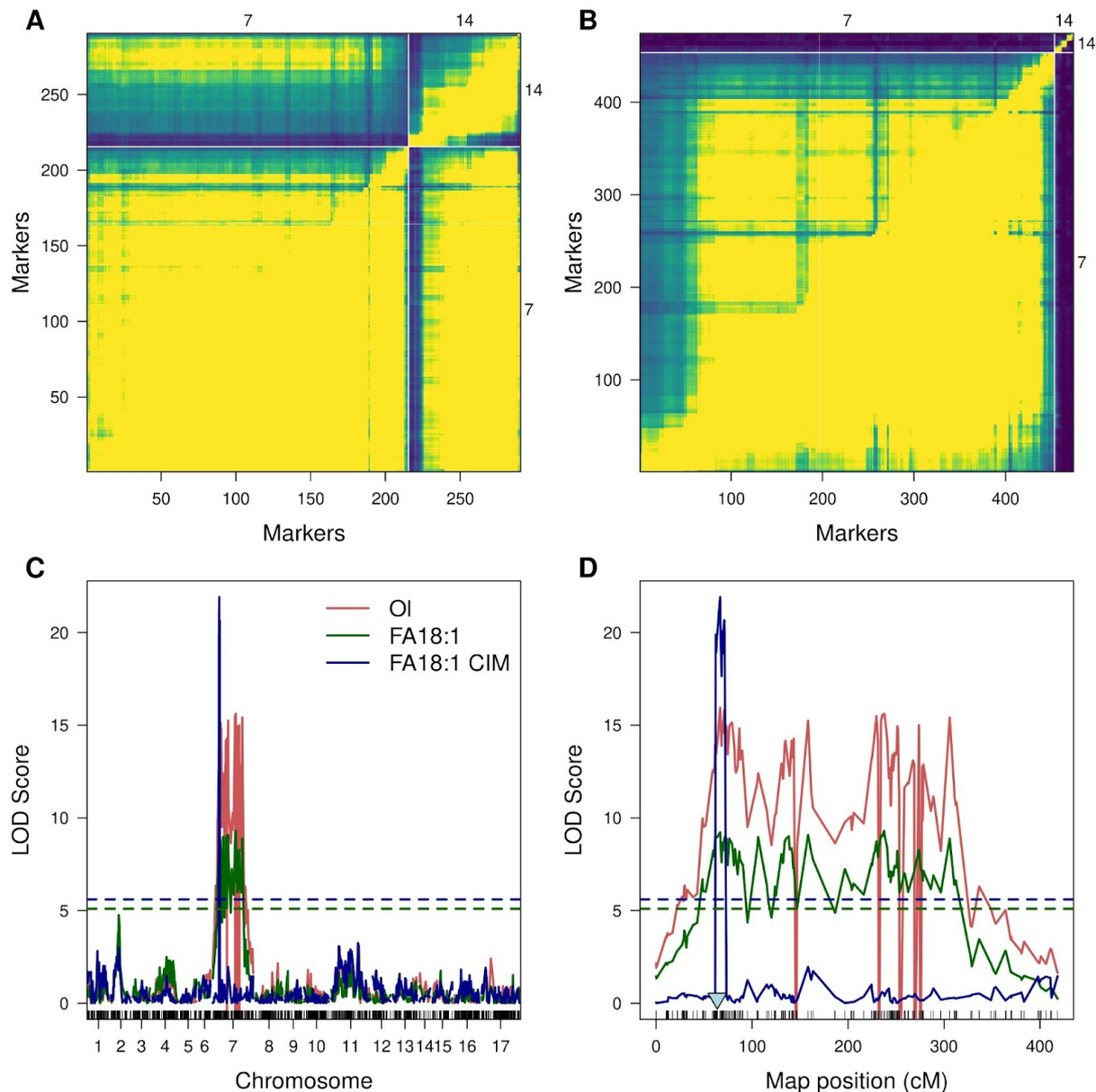


Figure 5.5.2 – Reassembling the chromosomes 7 and 14 for cross VK876xVK101 (Panels A and B) and updated mapping results (Panels C and D). The upper triangle of the matrices (panels A and B) corresponds to recombination fraction (RF), lower triangle corresponds to LOD scores. The blue color corresponds to high RF values and low LOD scores indicating poor linkage, while the yellow color corresponds to low RF values and high LOD indicating tight linkage between the pairs of markers.

Panel A reflects the RF and LOD scores for initial maps for chromosomes 7 and 14, panel B reflects RF and LOD scores for reassembled chromosomes 7 and 14. Red and green curves reflect the results of interval mapping of the binary *Ol*-associated trait and non-normally distributed oleic acid content, respectively. The blue curve reflects the results of composite interval mapping of oleic acid content. A light blue triangle indicates a marker located close to the *Ol* gene encoding FAD2-1.

In contrast, composite interval mapping detected only one locus located on chromosome 7 (Figure 5.5.2 D). Further analysis revealed the fact that the presence of loci located on chromosomes 7 and 14 could be explained by the tight linkage between markers from chromosome 7 and chromosome 14 (Figure 5.5.2 A, B). This could be the result of the potential translocation occurring between the markers from chromosome 7 and chromosome 14. Thus it was decided to reconstruct the genetic maps for markers from chromosomes 7 and 14 (Figure 5.5.2) without prior knowledge of the physical belonging of the markers to chromosomes. As a result, a significant amount of the genetic markers from chromosome 14 became a part of chromosome 7. Next, we performed the mapping with the re-assembled genetic map. As a result, we identified a single peak located on chromosome 7. Notably, the 1.5-LOD confidential interval obtained for significant markers identified by composite interval mapping from chromosome 7 spanning from 63.72 to 71.45 cM included one marker S14_78139380 which should be located on chromosome 14 based on the physical map (Table 5.5.1). Although it is derived from the 14th chromosome it was located 73 Mb away from the FAD2-1 gene which is quite distant even for F2 mapping that normally demonstrates low resolution.

Table 5.5.1 – Summary of genetic markers located within 1.5-LOD confidence interval resulted from composite interval mapping.

Marker	Cross	Chromosome	Position	Ol	FA18:1	FA18:1 CIM	PVE, %
S14_136819253	VK195xVK303	14	199.76	10.91	11.66	20.48	53.95
S14_137821336		14	202.86	11.91	12.01	22.27	56.37
S14_137795880		14	202.86	11.91	12.01	22.27	55.73
S14_137889873		14	203.49	11.85	11.82	21.96	56.07
S14_137951402		14	203.49	11.85	11.82	21.96	56.18
S14_137961667		14	203.49	11.85	11.82	21.96	56.41
S14_138978611		14	208.4	11.95	12.17	1.83	54.86
S14_78139380	VK876xVK101	7	63.72	13.95	8.96	19.88	46.99
S7_37821562		7	66.74	15.94	9.22	21.91	53.29
S7_39112905		7	66.74	15.94	9.22	21.91	50.46
S7_39776264		7	67.11	15.18	8.6	20.03	44.27
S7_39681220		7	67.11	15.18	8.6	20.03	47.39
S7_25411825		7	68.23	13.89	7.6	17.65	48.38
S7_25624783		7	68.23	13.89	7.6	17.65	46.14
S7_25717162		7	68.23	13.89	7.6	17.65	44.37
S7_25624784		7	68.23	13.89	7.6	17.65	46.14
S7_25664488		7	68.23	13.89	7.6	17.65	46.24
S7_25624756		7	68.23	13.89	7.6	17.65	46.14
S7_25421436		7	68.23	13.89	7.6	17.65	46.46
S7_25726978		7	68.23	13.89	7.6	17.65	44.37
S7_76273882		7	68.92	15.06	8.5	19.98	49.05
S7_25876036		7	68.92	15.06	8.5	19.98	48.62
S7_76319288		7	68.92	15.06	8.5	19.98	49.05
S7_25964564		7	68.92	15.06	8.5	19.98	48.83

S7_75968410		7	68.92	15.06	8.5	19.98	48.91
S7_76547907		7	68.92	15.06	8.5	19.98	49.11
S7_25961858		7	68.92	15.06	8.5	19.98	48.75
S7_25855919		7	68.92	15.06	8.5	19.98	48.62
S7_75140370		7	69.28	15.13	8.67	20.07	48.45
S7_72038884		7	69.28	15.13	8.67	20.07	51.02
S7_75826681		7	69.64	15.2	8.63	20.08	49.09
S7_75503792		7	69.83	14.59	8.23	18.88	46.9
S7_75503823		7	69.83	14.59	8.23	18.88	46.9
S7_75934390		7	70.02	15.2	8.63	20.08	49.01
S7_75892237		7	70.02	15.2	8.63	20.08	49.09
S7_75894251		7	70.02	15.2	8.63	20.08	49.09
S7_75535778		7	70.02	15.2	8.63	20.08	49.02
S7_76736113		7	71.1	15.83	8.78	20.66	50.33
S7_76758370		7	71.1	15.83	8.78	20.66	50.33
S7_76852814		7	71.45	15.2	8.96	19.73	48.72

The maximum proportion of variance explained by the genetic markers from the 1.5-LOD confidential interval was of 53.29% (Table 5.5.1, and Figure 5.5.3 B). Notably, unlike the significant markers from the VK195xVK303 the genetic markers from the VK876xVK101 demonstrated a dominant effect of the mutant *Ol* allele (Figure 5.5.3 B). This is in concordance with the identified segregation ratio of 3:1 reported for high-oleic and non-high oleic plants from cross VK876xVK101. The dominance of the genetic markers is also in agreement with the previously published results for the *Ol* gene (Lacombe et al., 2009; Lacombe & Bervillé, 2001; Schuppert et al., 2006).

It should be noted that the proportion of variance explained by the markers was similar compared to the previously published results that reported the range of 56 –

66% (Lacombe & Bervillé, 2001; Premnath et al., 2016). In addition, any additional loci that were previously described to affect the oleic acid content in addition to the major *Ol* gene (Premnath et al., 2016; F. Zhou et al., 2018) were not reported in the present study. This, first, could be related to the different genetic backgrounds of the lines from the present study and previously studied ones. Second, this could be due to the relatively low density of the genetic markers and sample size as well as the low resolution of F2 mapping populations.

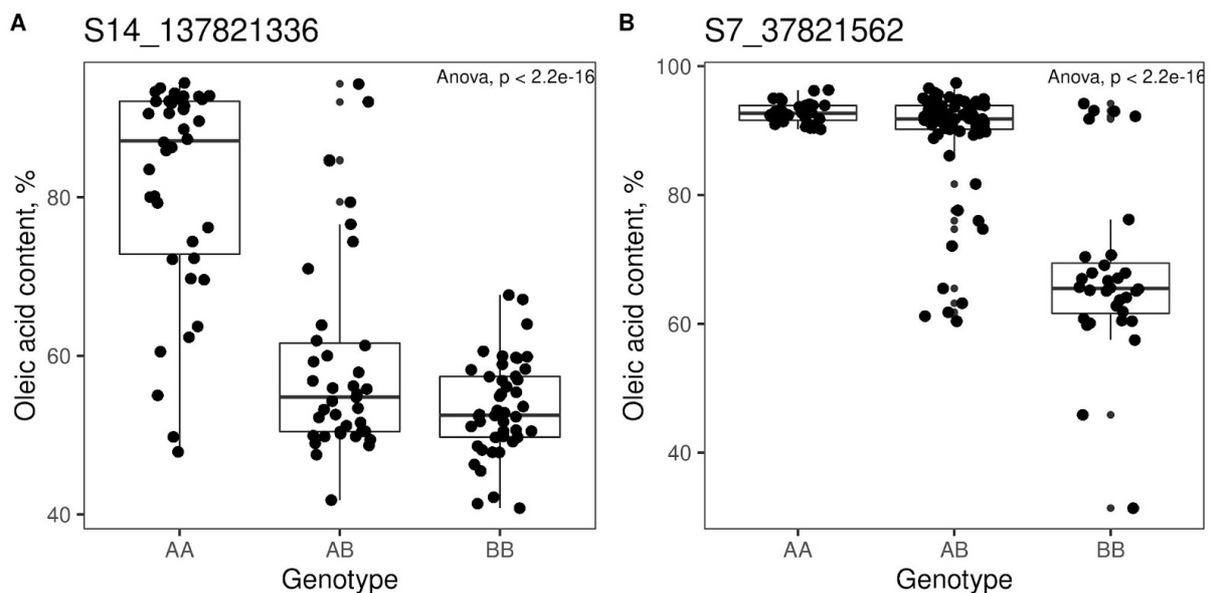


Figure 5.5.3 – Boxplots for genotype effects on oleic acid content. AA – genotype corresponds to maternal ones of a mutant line, BB – genotype wild ones derived from wild-type males. Panel A reflects the genotype effect of the most significant marker S14_137821336 for cross VK195xVK303. Panel B reflects the genotype effect of the most significant marker S7_37821562 for cross VK876xVK101.

The contradictory results of the unexpected effect of the recessive markers from cross VK195xVK303 laid the ground for deeper investigation of the *Ol* allele including sequencing of the targeted region and/or additional mapping experiments with the genetically contrasting parents. The potential translocation identified for cross VK876xVK101 could serve as a basis for the cytogenetic analysis of chromosomes 14 and 7 of lines VK876 and VK101. Additionally, a more precise

mapping of the major effect *Ol* gene is needed for this cross VK876xVK101. Although the potential genetic markers were identified for marker-assisted selection a deeper investigation of the loci potentially carrying the *Ol* gene is needed for both crosses.

5.6 Validation of the genetic markers

The genetic markers associated with the tocopherol composition were evaluated using Sanger sequencing. This was done to test the concordance between the alleles identified by GBS and Sanger sequencing being a "gold standard". To do so one marker for each of *Tph1* and *Tph2* and each of the two crosses (Table 5.6.1) was selected.

Table 5.6.1 – Summary of SNP verification.

Marker	Chromosome	Position (cM)	Population	Gene	LOD Score	Accuracy, %	Number matching genotypes
S1_55196434	1	88.71	VK195 X VK303	<i>Tph1</i>	20.18	94.12	32
S1_71748138	1	36.68	VK876 X VK101	<i>Tph1</i>	15.57	91.18	31
S8_23941299	8	79.34	VK876 X VK101	<i>Tph2</i>	24.66	85.29	29
S8_30578572	8	47.32	VK195 X VK303	<i>Tph2</i>	18.26	91.18	31

As a result, we tested four genetic markers. The main criteria for selection were the following: location of the marker within the 1.5-LOD confidence interval, unique sequence surrounding that makes it possible to construct primers for amplification of

a specific DNA region. Based on the above-mentioned criteria, the following markers were selected: S1_55196434 and S8_30578572 associated with *Tph1* and *Tph2* for the VK195xVK303 cross, respectively. For the VK876xVK101 cross, we selected S1_71748138 and S8_23941299 markers for *Tph1* and *Tph2*, respectively. As a result, four pairs of primers were constructed with the aim of amplification of four unique regions capturing the markers that underwent sequencing.

For verification of the markers, we selected 8 plants carrying the maternal (mutant) genotype, 8 plants carrying the paternal (wild-type) genotype and 8 ones carrying heterozygous genotypes based on the GBS data. Additionally, markers were tested on 5 female and 5 male parental plants. For this cohort, the allelic states of the markers were determined using Sanger sequencing and then compared to the allelic states obtained by GBS and calculated the proportion of matching genotypes. The genotypes obtained using the GBS approach that were used for genetic map construction and QTL mapping matched the genotypes obtained by Sanger sequencing with accuracy ranging from 85.29% to 94.12% (Table 5.6.1).

In addition to the verification of the allelic states with two different methods, an attempt was made to test whether the identified markers could be specific for non-relative lines. To test this, 20 genetically and phenotypically diverse lines were selected by collaborators from VNIIMK (Figure 5.6.1). These lines were sequenced in 5 replicates and phenotypically characterized in terms of the presence/absence of *Tph1*, *Tph2* (four phenotype classes) and *Ol* (high-oleic and non-high oleic) alleles of genes based on the tocopherol composition and oleic acid content previously reported for these lines.

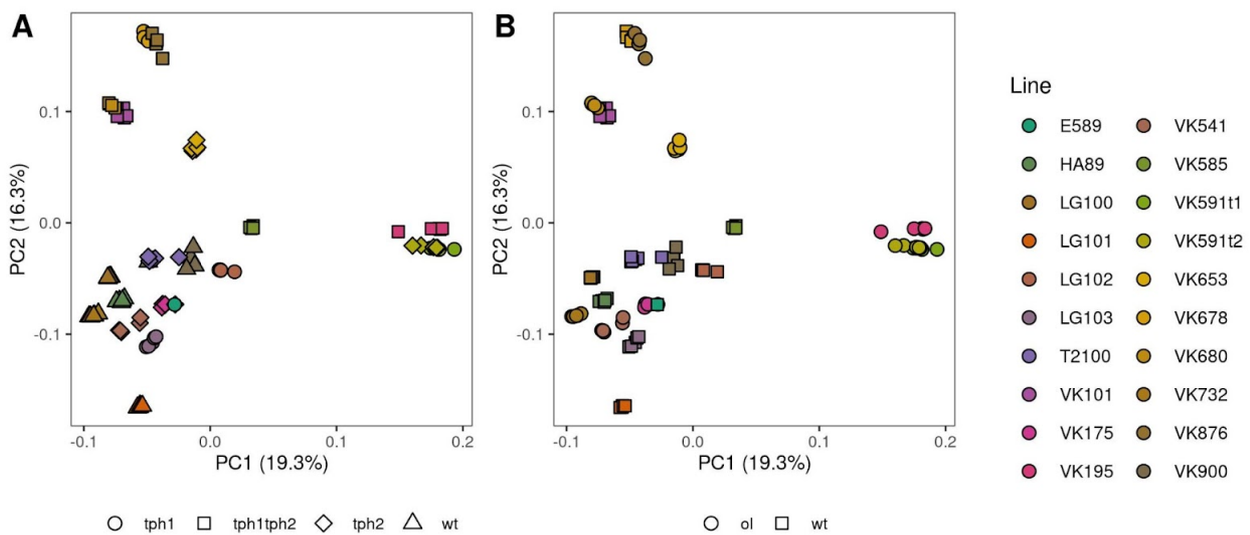


Figure 5.6.1 – Principal component analysis of the population structure of the verification sample. Color corresponds to the sunflower line. The shape of the figures reflects the phenotype associated with either tocopherol composition (Panel A) or oleic acid content (Panel B). The numbers in brackets correspond to the proportion of genotype variance explained by the principal component.

After the sequencing data was obtained, joint SNP calling with experimental cross samples was performed with the aim of finding genetic polymorphisms common in verification sample and experimental crosses. Next, 10 most significant markers were selected for each of the trait/population combinations. Next, *Ol*-, *Tph1*- and *Tph2*-associated phenotypes of the verification samples were predicted based on the allelic states of the significant markers identified either with VK195xVK303 or VK876xVK101 populations (Figure 5.6.2). It was demonstrated that the *Tph2*-associated phenotypes were most accurately predicted with the genetic markers identified with both populations. In particular genetic markers S8_30578572 mapped using VK195xVK303 cross and S8_24695806 mapped using VK876xVK101 cross demonstrated accuracy of 88.8% and 87.5%, respectively, being genotyped in 90% and 80% of lines from verification sample.

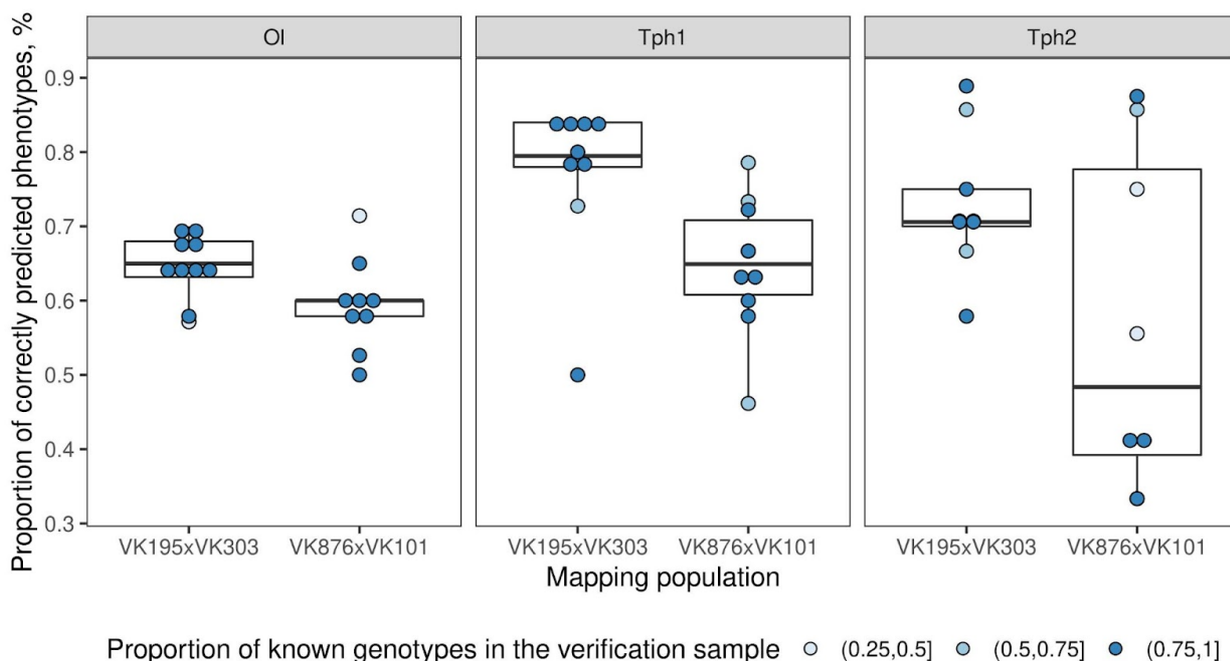


Figure 5.6.2 – Accuracy of the genetic markers to predict *Ol*, *Tph1* and *Tph2*-associated phenotypes (indicated by facets). Each dot corresponds to the marker. The color of the dot represents the proportion of known genotypes that were genotyped among individuals from the verification sample. The abscissa axis indicates the population for which the genetic marker was tested. The ordinate axis reflects the proportion of the correctly predicted phenotypes associated with *Ol*, *Tph1* and *Tph2*.

Such a high proportion of correctly predicted phenotypes and genotyping level could be explained by the physical proximity of these markers to the causal gene located on chromosome 8. For *Tph1* genetic markers S1_56474204, S1_55157943, S1_55196434 and S1_54474863 obtained by analyzing VK195xVK303 cross demonstrated the accuracy of 84% being identified in the more than 90% of verification lines. From cross VK876xVK101 only one genetic marker S1_71748138 predicted a *Tph1*-associated phenotype with an accuracy of 78.8%, the accuracy of the other markers was lower than 75%. For high oleic traits associated with *Ol*, the accuracy of the genetic markers was lower than 75%. In particular marker S7_76852814 from the cross, VK876xVK101 demonstrated a maximum accuracy of

71% and marker S14_136584727 from cross VK195xVK303 demonstrated a maximum accuracy of 70%. Lower predicting accuracy could be related to the fact that these markers were located far away from the causal gene FAD2-1, additionally for VK195xVK303 markers demonstrated unexpected recessive nature, while for population VK876xVK101 genetic markers were linked to chromosome 7, which was not previously reported to carry major effect gene encoding FAD2-1.

5.7 Conclusions

In the present part of the study the mapping of genetic markers associated with the oil quality was performed, namely tocopherol composition and oleic acid content. Two F2 populations derived from parental lines with contrast tocopherol composition and oleic acid content were analyzed. A GBS approach followed by SNP calling, imputation and genetic map construction was applied with the aim of mapping these traits. Here quantitative and qualitative mapping approaches were applied for both traits as it was assumed that in both crosses the two major effect genes *Tph1* and *Tph2* for tocopherol composition and *Ol* for oleic acid content control the trait expression. This assumption is on the one hand supported by the fact that the phenotypic segregation ratios of 9:3:3:1 and 3:1 for tocopherol composition and oleic acid content were observed, indicating a mono/di-genic mode of inheritance for these traits. On the other hand, quantitative mapping of these traits did not reveal any additional loci that control these traits.

Notably, in the previous studies, additional loci controlling tocopherol composition were identified (García-Moreno et al., 2012b; Hass et al., 2006). For oleic acid content, minor effect loci controlling this trait were discussed in addition to the major effect gene (Premnath et al., 2016; F. Zhou et al., 2018). The fact that here no additional loci were identified could be explained by first that here we used a relatively small cohort, second by the different genetic backgrounds of the lines from the present and past studies.

Due to the applied aim of this study associated with the scanning for genetic markers of oil quality, first a GBS approach was validated using the Sanger sequencing, to prove the concordance of the markers' alleles by two independent approaches. Second, genetic markers were tested for the ability to predict the phenotypes associated with major effect genes *Tph1*, *Tph2* and *Ol* in independent plant samples. These results demonstrated that *Tph1*- and *Tph2*-associated markers were identified in a large proportion of the test sample including unrelated lines (> 90%) and predicted associated phenotypes with high accuracy (> 84%). Thus, we selected a set of genetic markers that will be used to develop a marker-assisted selection approach for sunflower oil improvement in the context of tocopherol composition.

Regarding the oleic acid content, the genetic markers demonstrated lower parameters in the context of accuracy and the presence in the test sample. This could be due to the fact that the genetic markers associated with oleic acid content for population VK195xVK303 were recessive, which on the one hand could be related to the genetic background and on the other a potential existence of an alternative allele of *Ol* behaving as recessive ones. For cross VK876xVK101 identified markers were dominant, however due to possible translocation between chromosomes 14 and 7, most of the genetic markers were attributed to chromosome 7 which was not previously reported to carry genes associated with oleic acid content and *Ol* gene in particular. It also should be noted that in both crosses the genetic markers associated with *Ol* were located more than 13 megabases away from the causal gene FAD2-1 based on the physical map information which in turn may indicate poor linkage with this gene. All these discordances with the initially set hypothesis could be set as goals for future studies for a better understanding of *Ol* gene inheritance in the sunflower in lines from the VNIIMK collection.

Chapter 6. Association mapping of seed morphology traits in sunflower

6.1 Introduction

One of the key phenotypes of the sunflower that is of high interest to breeders and industry is seed-related traits – seed size, husk size and seed to husk ratio. Seed and husk size are of particular interest in the development of large-seeded confectionary sunflowers, additionally (Lukomets et al., 2021). One of the methods used to estimate seed size is a measurement of the weight of thousand seeds. This technique was widely applied to oilseed crops (Khan et al., 2019; Souza et al., 2016), including the sunflower (Radic et al., 2013) as the seed length and width significantly correlate with the thousand seed weight (Gjorgjieva et al., 2015). However, the main disadvantage of this method is that it does not provide information on the ratio of seed and husk area, being an important parameter of whether the photoassimilates are rationally utilized by plants in terms of increasing the yield output. One of the ways to estimate seed size, husk size and seed to husk ratio in a non-invasive manner is to perform x-ray radiography methods followed by image analysis (Arkhipov et al., 2019). Previously the X-ray computer methods were mostly applied to assess morphological aspects of the seeds (Rocha et al., 2014), quality including viable seeds, empty seeds and seeds damaged by the pathogen (Dumont et al., 2015), seed viability (Al-Turki & Baskin, 2017). In the present part of the study genetic mapping of seed-related traits collected by means of the X-ray computer methods by our collaborators from St. Petersburg Electrotechnical University LETI was performed. Traits that were collected included husk area, seed area as well as a seed-to-husk ratio. These characteristics were collected for the genetically and phenotypically diverse collection of the sunflower which included a total of 601 accessions from VIR and VNIIMK institutions and the Agroplasma breeding company. The part of the results related to the genetic diversity assessment of the 601 lines in the following chapter was published in the BMC Genomics journal (Chernova et al., 2021).

6.2 Collection of phenotypes

To collect the information on the seed size, husk size and seed to husk ratio the seeds were placed on the plate (Figure 6.2.1) and next scanned using PRDU-02 setup followed by analysis using SeedRentgen. As a result, mean values were collected for the seed area and husk area (Table 6.2.1).

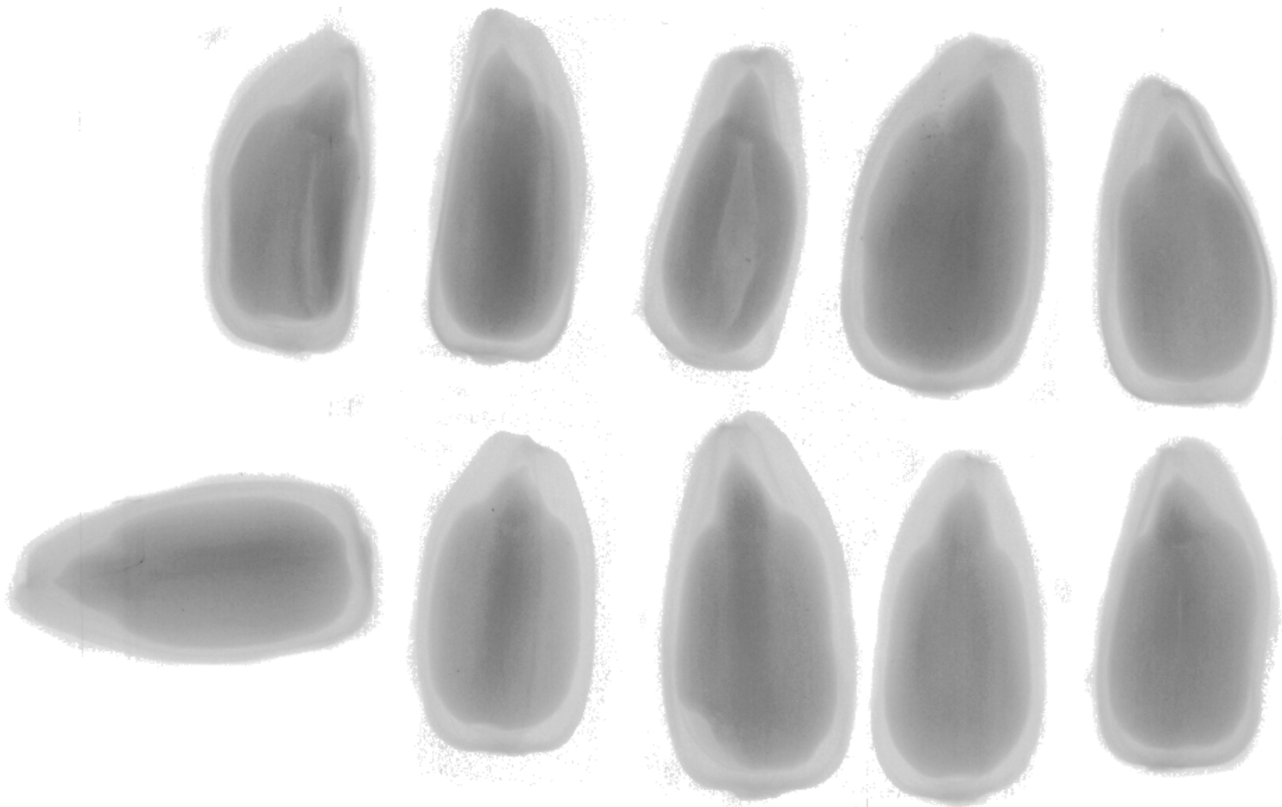


Figure 6.2.1 – Example of X-ray radiograph with inverted color. Each radiograph was made for 10 seeds of accession. Original pictures are with a black background and white seeds. The size of the pixel corresponds to 0.1 mm^2 .

Table 6.2.1 – Summary table on seed-related traits. Mean values \pm standard deviation are shown 

Collection	Husk area, mm ²	Seed area, mm ²	Husk to seed ratio
VIR	15.01 ± 4.53	7.51 ± 2.01	51.15 ± 7.01
VNIIMK	39.88 ± 8.11	19.28 ± 3.85	49.11 ± 8.34
AGROPLASMA	37.06 ± 11.68	20.56 ± 5.21	57.10 ± 7.75
Mean	28.54 ± 14.21	14.54±7.10	51.92 ± 8.23

Analysis of variance revealed a significant difference of seed and husk area between the collections (Figure 6.2.2). Namely, husk was on average 2.65 times smaller in VIR accessions when comparing VIR and VNIIMK collections (Mann–Whitney U test p-value < 2.2e-16) collections and on average 2.46 times smaller when comparing VIR and AGROPLASMA collections (Mann–Whitney U test p-value < 2.2e-16). The difference between AGROPLASMA and VNIIMK collection was far less prominent (1.07 times), but still significant (Mann–Whitney U test p-value = 0.012). The same pattern applied to the seed area: on average VIR accessions were 2.56 times smaller than VNIIMK accessions (Wilcoxon test p-value < 2.2e-16) and 2.73 times smaller than AGROPLASMA accessions. The difference between VNIIMK and AGROPLASMA was marginally significant (Mann–Whitney U test p-value = 0.041) and was 1.07 times.

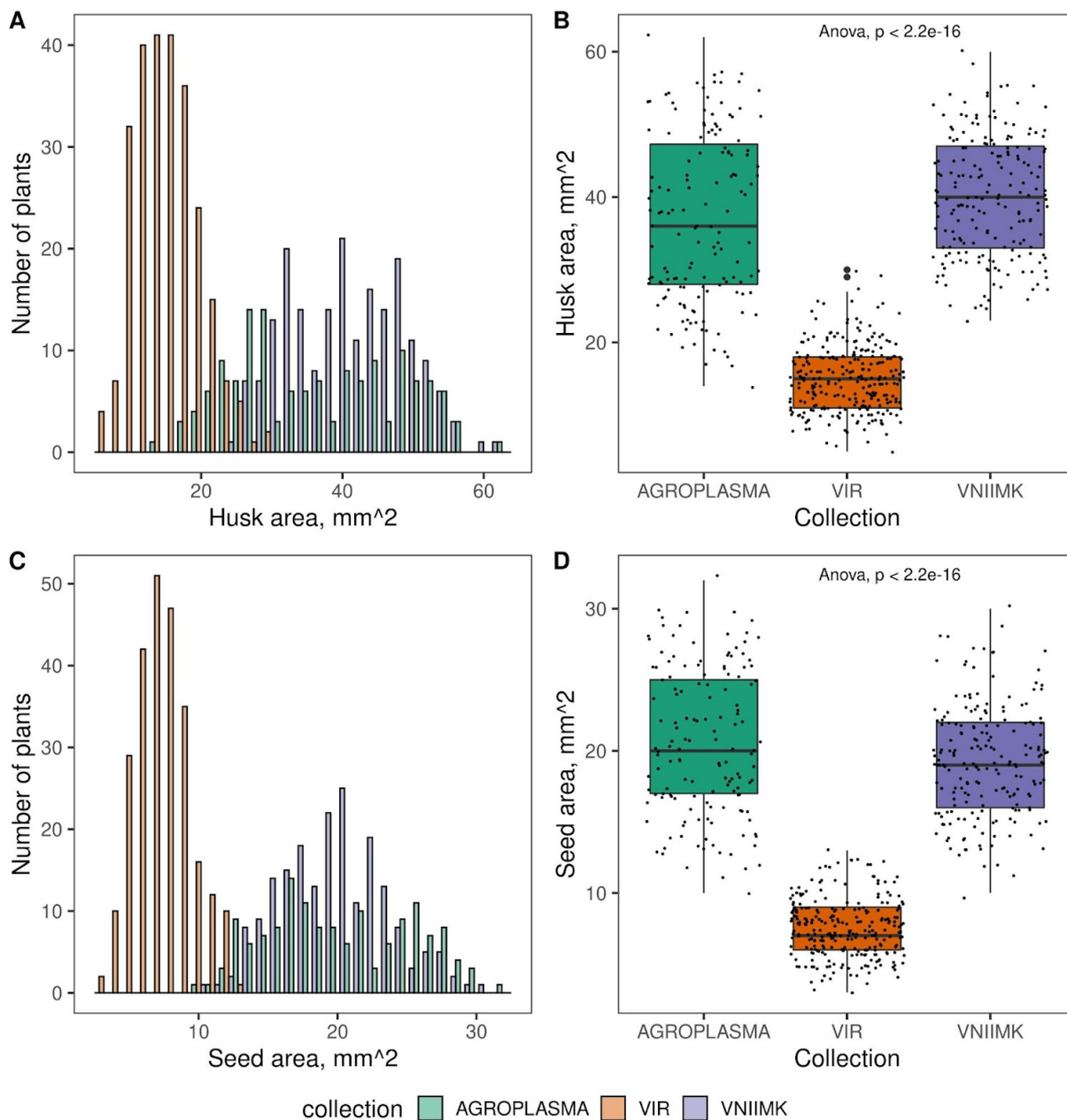


Figure 6.2.2 – Distribution of the husk (panel A and B) and seed (panels C and D) area between the collections. The color of the bars on histogram and box plots corresponds to the collection of the seed. Each dot corresponds to the mean value obtained for 10 seeds from the X-ray radiograph.

Next, a relationship between the husk and seed area was estimated by assessing the Pearson correlation coefficient. For the whole set of observations, the correlation between husk and seed area was significant ($R = 0.93$, $p\text{-value} < 2.2e-16$). However, it differed while being assessed separately for three collections (Figure 6.2.3)

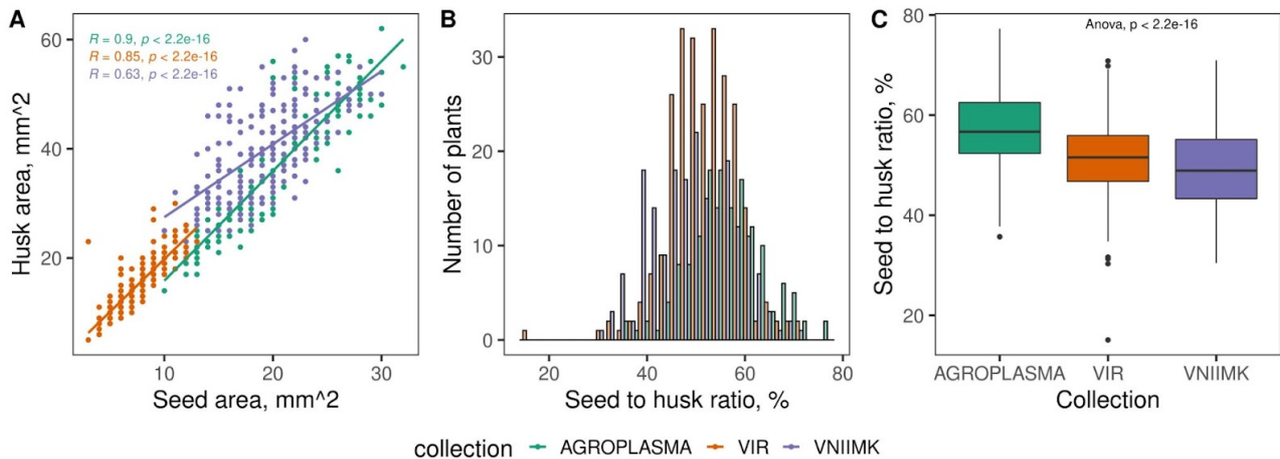


Figure 6.2.3 – Correlation between seed and husk area (panel A), distribution of the seed to husk ratio among collections. The color of regression lines, histogram bars and box plots correspond to the collection of the seed.

Additionally, the seed to husk ratio differed between collections (ANOVA p-value < 2.2e-16). Here an Agroplasma collection demonstrated the highest value of 57.10 which was significantly larger than the VIR sample by 1.11 times (Mann–Whitney U test p-value = 2.428e-13) and VNIIMK samples by 1.16 times (Mann–Whitney U test p-value = 4.651e-16).

Such differences in seed and husk sizes could be explained by the fact that the VIR collection mostly consists of historical samples, while VNIIMK and AGROPLASMA collections include economically valuable accessions used to produce commercial hybrids and lines for which the size of the seed is one of the key traits to be improved. Thus, the differences are explained by the artificial selection pressure different for the studied collections. The same could be applied to the difference in the seed to husk ratio the highest value was specific for the AGROPLASMA collection which also speaks in favor of the particular interest of increasing the seed area compared to the husk area so that the volume of the kernel was high. The performed analysis established that the collection of the accession is a

key factor affecting the seed-related traits. Thus, this factor is used as a covariate for subsequent association mapping.

6.3 Assessment of genetic diversity

To describe genetic diversity and perform mapping of the agronomically important traits, 15068 SNPs were obtained for 601 accessions from three collections by means of GBS followed by GATK pipeline analysis. As it was the first study that includes a substantial set of the genetically diverse sunflower accession collection, a characterization of the population structure was performed (Figure 6.3.1).

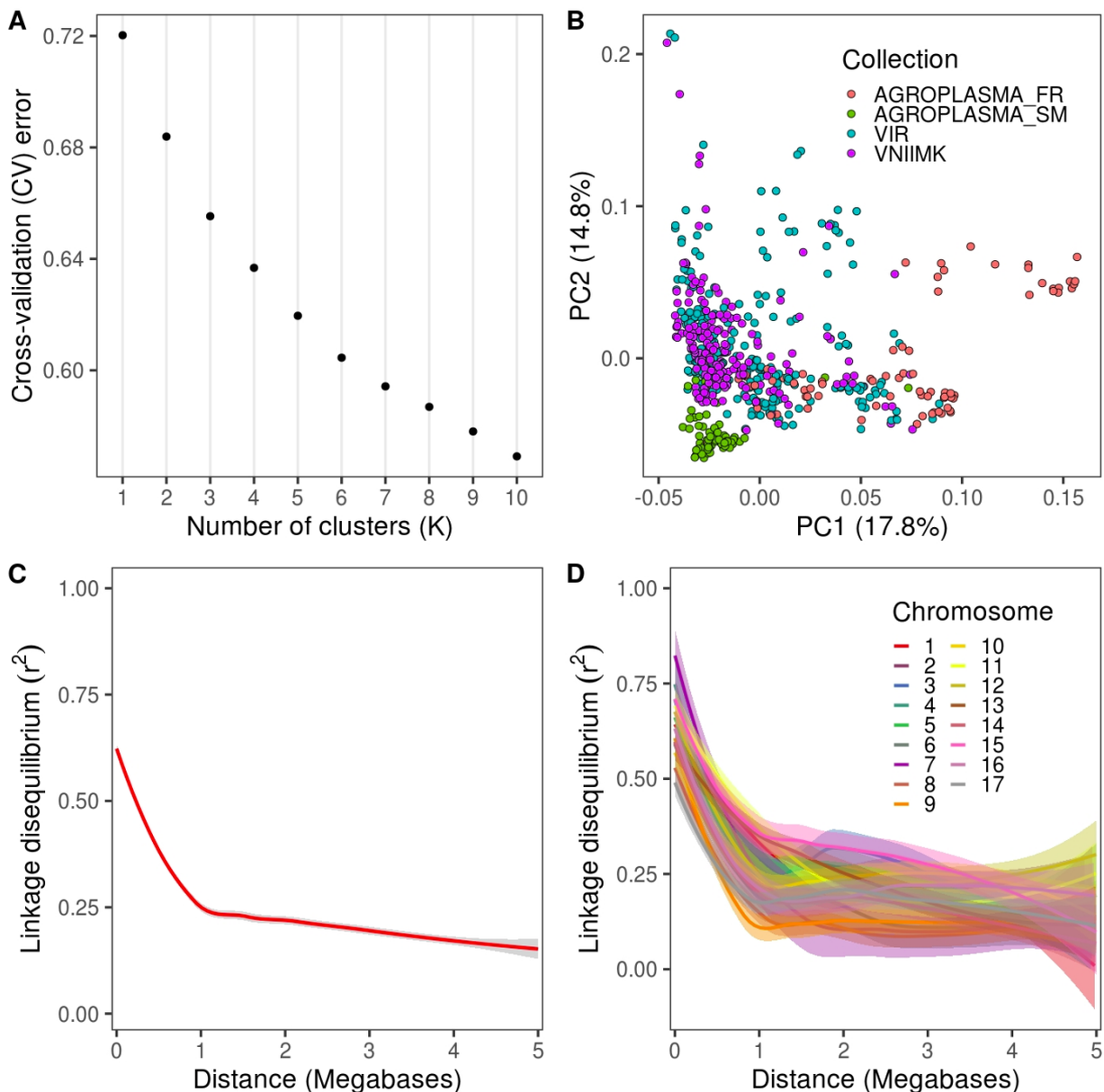


Figure 6.3.1 – Population structure of the 601 sunflower accession from VIR, VNIIMK and Agroplasma collections. Cross-validation error values for K ranging from 1 to 10 estimated by ADMIXTURE software (panel A). Visualization of first two principal components of PCA (panel B). Linkage disequilibrium decay across the whole genome (panel C) and separate chromosomes (panel D).

An ADMIXTURE and PCA analysis did not reveal any clear population structure, neither between collections, nor between phenotypically diverse lines. However, slight differences were identified for fertility restorer and fertility maintainer lines of the Agroplasma collection (Figure 6.3.1). Next, we assessed a Linkage disequilibrium decay for the whole genome as well as for the separate chromosomes. While the average genotype correlation (r^2) dropped to half of its maximum value at 0.7 Mb, linkage disequilibrium (LD) decay varied among the 17 chromosomes.

As the present study was the first one devoted to the genotyping of a significant amount of the accession from Russian collections the genetic differences with international collections that included wild representatives of the *Helianthus* genus, cultivated and landrace accessions were compared. To do so a principal component was performed on the 2345 SNPs shared between the collection of more than 1000 accessions (Hübner et al., 2019) for which an unfiltered VCF was provided by colleagues from the Biodiversity Research Centre of the University of British Columbia.

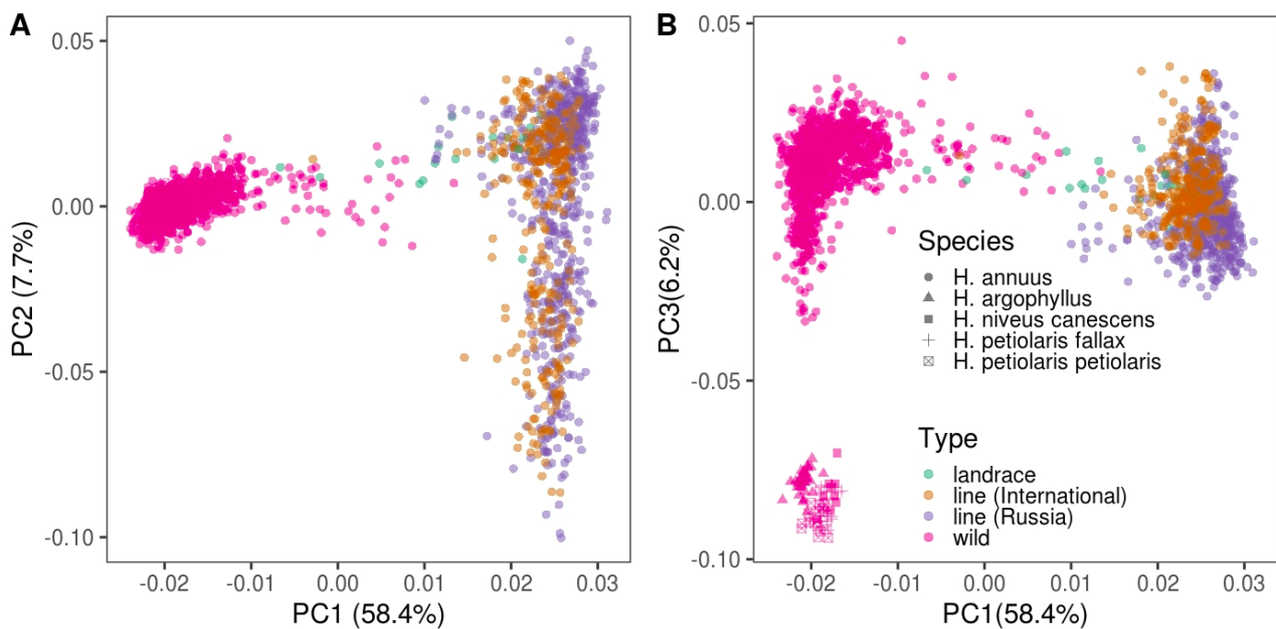


Figure 6.3.2 – Joint principal component analysis of sunflower accessions genotyped in this study and Hübner (2019) based on 2345 shared SNPs. The first and the second (A) or the first and the third (B) PCs are shown. Each dot corresponds to a plant accession. Shapes correspond to species. The origin (wild/line/landrace) is indicated by color.

The principal component analysis revealed a clear separation of wild and cultivated sunflowers by the first principal component explaining 58.4% of genotype variance. The first and third principal components revealed three clusters that consisted of wild *Helianthus annuus*, cultivated *Helianthus annuus*, and the third cluster was represented by other wild representatives of the *Helianthus* genus (Figure 6.3.2 B). The first three PCs demonstrated that the genetic diversities of the cultivated Russian and International sunflower accessions are matching.

6.4 Association mapping of seed-related traits and scanning for the candidate genes

As the information on the genotypes and phenotypes were collected, an association mapping of husk and seed areas as well as the seed-to-husk ratio was performed. To do so, a compressed mixed linear model approach was applied within the TASSEL software. For each trait, the first five principal components as well as

the kinship matrix were added as confounding factors. Additionally, information on the collection was also added as a covariate as it significantly affected the distribution of all three traits.

As a result, only one SNP located on chromosome 10 associated with the husk area overcame the Bonferroni corrected threshold. Thus, a softer threshold of $p=0.0001$ was selected. As a result, we identified additional husk area-associated SNPs located on chromosomes 4, 9 and 17 (Figure 6.4.1). The proportion of the trait variance explained by these SNPs varied from 3.65 to 6.49 percent (Table 6.4.1).

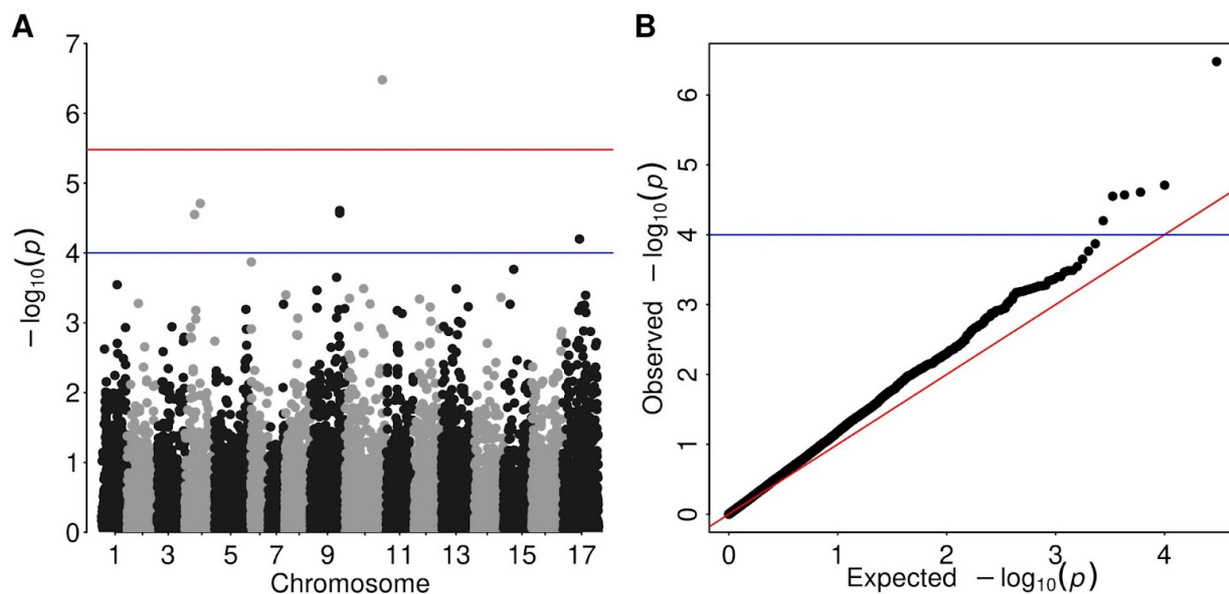


Figure 6.4.1 – Manhattan plot for association mapping of husk area results (panel A) and corresponding Q-Q plot (panel B). each dot corresponds to SNP. Black dots on panel A correspond to SNPs from odd chromosomes, gray dots correspond to SNPs from even chromosomes. The red horizontal line on panel A corresponds to the Bonferroni corrected p-value of 0.05. Blue horizontal lines on panels A and B correspond to a p-value threshold equal to 0.0001.

For the seed area, none of the SNPs overcame the Bonferroni multiple testing correction threshold. However, eight SNPs demonstrated a p-value below the softer

threshold of $p=0.0001$. These SNPs were located on chromosomes 2, 6, 9, 10, 11, 13 and 14 (Figure 6.4.2). The proportion of the trait variance explained by these SNPs varied from 3.82 to 5.52 percent.

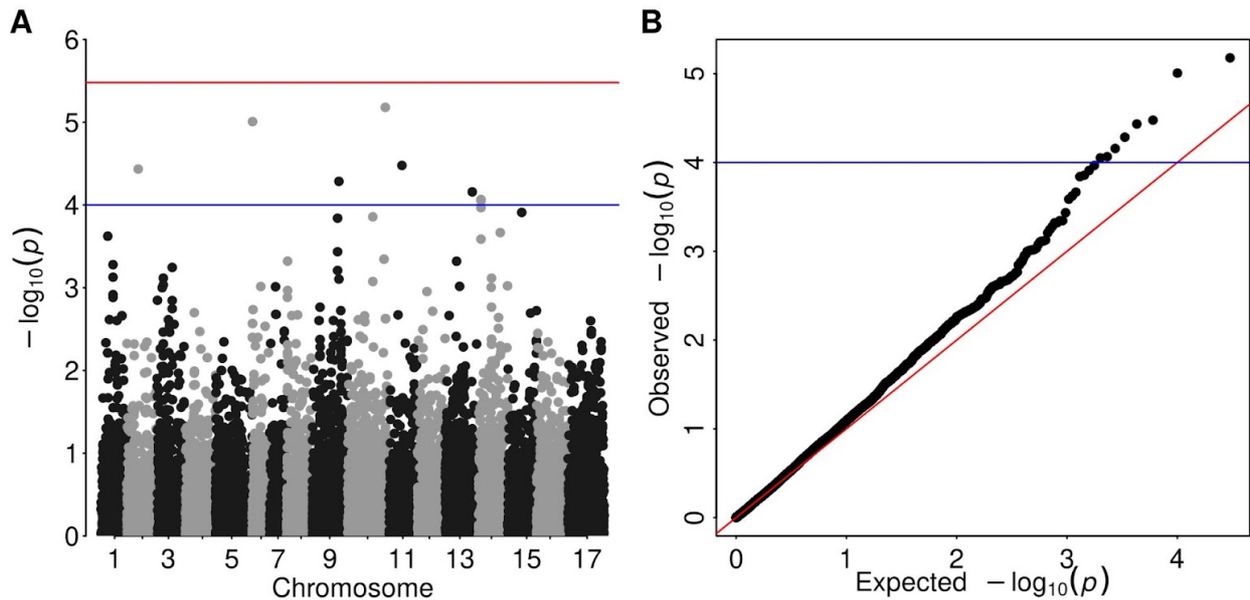


Figure 6.4.2 – Manhattan plot for association mapping of seed area results (panel A) and corresponding Q-Q plot (panel B). Each dot corresponds to the SNP. Black dots on panel A correspond to SNPs from odd chromosomes, gray dots correspond to SNPs from even chromosomes. The red horizontal line on panel A corresponds to the Bonferroni corrected p-value of 0.05. Blue horizontal lines on panels A and B correspond to a p-value threshold equal to 0.0001.

Finally, for the seed to husk ratio trait only one chromosome, chromosome 10, carried SNPs associated with the trait under the softer statistical threshold (Figure 6.4.3). The corresponding SNPs explained from 3.39 to 5.05 percent of trait phenotypic variation (Table 6.4.1).

Thus, the low amount of the phenotypic variance explained by the markers was reported for all of the studied traits in the present study. The previous studies focusing on the association mapping of the seed size-related traits (seed length, seed width, seed area, kernel length, kernel length, seed weight) identified loci controlling

these traits that explained a higher proportion of phenotypic variance compared to the present study.

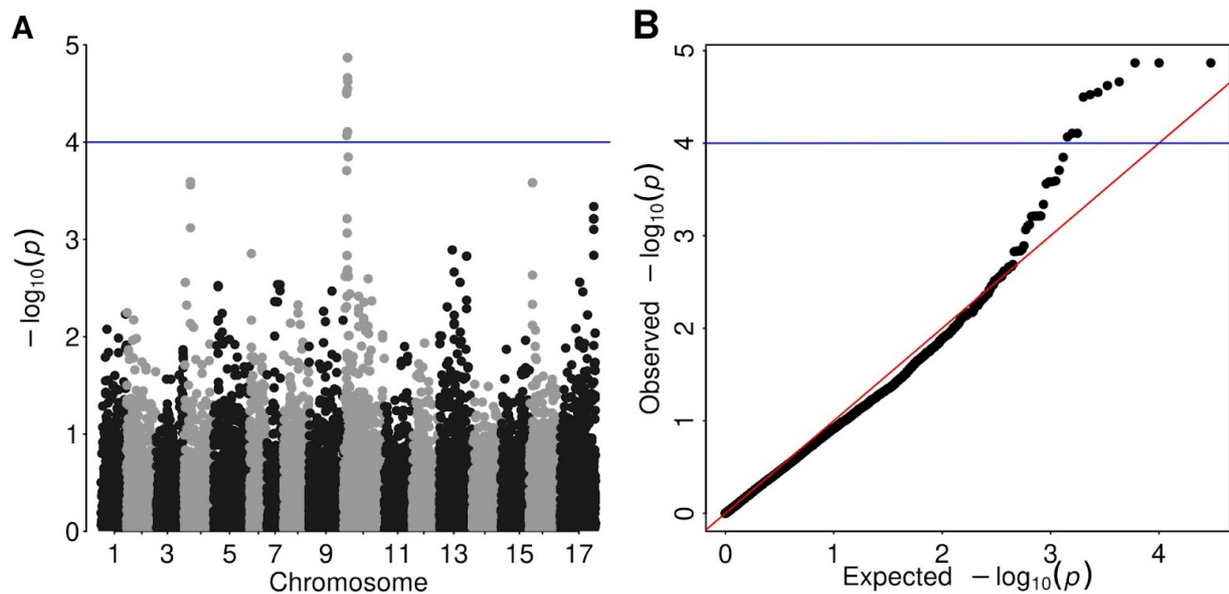


Figure 6.4.3 – Manhattan plot for association mapping of seed to husk ratio (panel A) and corresponding Q-Q plot (panel B). Each dot corresponds to the SNP. Black dots on panel A correspond to SNPs from odd chromosomes, gray dots correspond to SNPs from even chromosomes. Blue horizontal lines on panels A and B correspond to a p-value threshold equal to 0.0001.

A QTL mapping performed on the F2 mapping populations revealed seven genetic markers associated with 100-seed weight, seed length and seed width jointly explaining 56, 42 and 65 percent of phenotypic variance, respectively (Tang, Leon, et al., 2006). Another QTL study identified loci associated with the seed length, seed width, kernel length and kernel width explaining up to 14.8, 10.7, 13.5 and 23.2 percent of phenotypic variance, respectively. Associated loci were identified on chromosomes 1, 4, 7, 10, 13, 16 and 17 (Yue et al., 2009). A GWAS mapping of seed length, seed width and seed area based on sunflower diversity panels identified loci explaining up to 15, 85 and 47 percent of phenotypic variance, respectively. Loci associated with these traits were mapped on chromosomes 10, 13 and 17 (Reinert et

al., 2020). Additionally, it was previously demonstrated that the seed weight, which correlates well with the seed size, demonstrates a high level of heritability ranging from 79 to 89 percent in sunflower (Abdelsatar et al., 2020; Baraiya & Patel, 2018; A. Khan, 2001). Despite the fact that previously identified loci explained more variance, compared to the present study, they still explained less variation than expected considering the high rate of seed-related traits heritability. This discrepancy could be referred to as the missing heritability phenomenon discussed for plants previously (Brachi et al., 2011). Further, incomplete agreement between our current results and published studies, as well as among reported loci, indicate the need for further large-scale genetic studies of the seed-related phenotypes.

Table 6.4.1 – Summary of genetic markers associated with seed-related traits.

Marker	Trait	Chromosome	Position	p-value	PVE, %
S10_18982408	seed_husk_ratio	10	18982408	2.99E-05	4.33
S10_18982669	seed_husk_ratio	10	18982669	3.17E-05	4.21
S10_19187693	seed_husk_ratio	10	19187693	8.55E-05	3.39
S10_24409435	seed_husk_ratio	10	24409435	1.35E-05	4.85
S10_24409447	seed_husk_ratio	10	24409447	1.35E-05	4.85
S10_24409448	seed_husk_ratio	10	24409448	1.35E-05	4.85
S10_24409452	seed_husk_ratio	10	24409452	2.17E-05	5.05
S10_24409622	seed_husk_ratio	10	24409622	7.82E-05	4.32
S10_24409623	seed_husk_ratio	10	24409623	7.82E-05	4.32
S10_24409658	seed_husk_ratio	10	24409658	2.82E-05	4.57
S10_25468639	seed_husk_ratio	10	25468639	2.38E-05	4.56
S04_56847151	husk	4	56847151	2.81E-05	4.22
S04_91194852	husk	4	91194852	1.95E-05	3.65

S09_175848372	husk	9	175848372	2.46E-05	3.68
S09_175848377	husk	9	175848377	2.69E-05	3.66
S10_222897158	husk	10	222897158	3.32E-07	6.49
S17_99160416	husk	17	99160416	6.31E-05	3.78
S02_67897490	seed	2	67897490	3.68E-05	4.22
S06_1852907	seed	6	1852907	9.83E-06	4.95
S09_157078177	seed	9	157078177	5.18E-05	4.15
S10_222897158	seed	10	222897158	6.61E-06	5.52
S11_76005476	seed	11	76005476	3.33E-05	4.51
S13_159859510	seed	13	159859510	6.95E-05	4.09
S14_14459174	seed	14	14459174	8.86E-05	3.82
S14_14459481	seed	14	14459481	8.59E-05	3.84

We next performed a scanning for the causal genes within the associated loci. To do so, we first selected genes located within 100kb interval upstream and downstream the associated markers ; as SNP pairs located within this frame demonstrated a high level ($r^2 > 0.5$) of linkage (Figure 6.3.1). As a result, more than 100 genes located within the selected frame were identified. Most of the genes encoded proteins of unknown functions, however some of the genes included meaningful annotation (Table 6.4.2) and encoded proteins that could be potentially involved in the seed size-related parameters.

Table 6.4.2 Genes with annotation located within 100kb frame upstream and downstream significant SNPs.

Trait	SNP	GeneID	Product	Annotation	Distance
husk	S04_91194852	HannXRQ_Chr04g0109941	Putative cysteine-rich secretory protein allergen V5/Tpx-1-related	intergenic_region	0

husk	S09_175 848377	ATMLH1	NA	downstream_gene _variant	23392
husk	S09_175 848377	HannXRQ_Chr0 9g0266421	Putative UDP-glucuronosyl/UDP-glucosyltransferase	missense_variant	0
husk	S09_175 848377	HannXRQ_Chr0 9g0266441	Putative cytochrome P450	downstream_gene _variant	99156
husk	S10_222 897158	HannXRQ_Chr1 0g0312591	Putative cytochrome P450	upstream_gene_v ariant	22240
husk	S10_222 897158	HannXRQ_Chr1 0g0312611	Putative transcription factor GRAS Scarecrow-like 29	downstream_gene _variant	37826
husk	S10_222 897158	HannXRQ_Chr1 0g0312621	Putative germin	downstream_gene _variant	90730
husk	S17_991 60416	HannXRQ_Chr1 7g0555021	Putative protein kinase-like domain, Leucine-rich repeat-containing N-terminal	downstream_gene _variant	84262
husk	S17_991 60416	HannXRQ_Chr1 7g0555031	Probable S-adenosyl-L-methionine-dependent methyltransferases superfamily protein	intergenic_region	0
husk	S17_991 60416	HannXRQ_Chr1 7g0555031	Probable S-adenosyl-L-methionine-dependent methyltransferases superfamily protein	upstream_gene_v ariant	14634
seed	S06_185 2907	HannXRQ_Chr0 6g0164571	Probable translation elongation factor EF1B	downstream_gene _variant	47433
seed	S06_185 2907	HannXRQ_Chr0 6g0164581	Probable ATP synthase D chain mitochondrial	downstream_gene _variant	40865
seed	S06_185 2907	HannXRQ_Chr0 6g0164591	Putative tetraspanin/Peripherin	upstream_gene_v ariant	34142
seed	S06_185 2907	HannXRQ_Chr0 6g0164601	Putative proton-dependent oligopeptide transporter family Major facilitator superfamily domain	downstream_gene _variant	8529
seed	S06_185 2907	HannXRQ_Chr0 6g0164611	Putative proton-dependent oligopeptide transporter family Major facilitator superfamily domain	synonymous_vari ant	0
seed	S09_157 078177	HannXRQ_Chr0 9g0261661	Putative tetratricopeptide-like helical domain	upstream_gene_v ariant	33065
seed	S09_157 078177	HannXRQ_Chr0 9g0261671	Probable quinone reductase family protein	downstream_gene _variant	23147
seed	S09_157 078177	HannXRQ_Chr0 9g0261681	Putative calcium/proton exchanger	upstream_gene_v ariant	19803
seed	S10_222 897158	HannXRQ_Chr1 0g0312581	Putative dnaJ domain	downstream_gene _variant	40006
seed	S10_222 897158	HannXRQ_Chr1 0g0312591	Putative cytochrome P450	upstream_gene_v ariant	22240
seed	S10_222 897158	HannXRQ_Chr1 0g0312621	Putative germin	downstream_gene _variant	90730
seed	S11_760 05476	HannXRQ_Chr1 1g0335701	Probable SAUR-like auxin-responsive protein family	upstream_gene_v ariant	3130

seed	S11_760 05476	HannXRQ_Chr1 1g0335721	Putative meiotic nuclear division protein 1	downstream_gene _variant	53138
seed	S13_159 859510	HannXRQ_Chr1 3g0416301	Putative cytochrome P450	downstream_gene _variant	52185
seed	S13_159 859510	HannXRQ_Chr1 3g0416311	Probable peptidase M1 family protein	upstream_gene_v ariant	76515
seed	S14_144 59481	BSK1	BR-signaling kinase 1	upstream_gene_v ariant	63202
seed_hus k_ratio	S10_189 82669	HannXRQ_Chr1 0g0281741	Putative nucleotide-binding alpha-beta plait domain	intergenic_region	0
seed_hus k_ratio	S10_244 09435	HannXRQ_Chr1 0g0282351	Putative ankyrin repeat-containing domain Gag-polypeptide of LTR copia-type	downstream_gene _variant	93932
seed_hus k_ratio	S10_244 09435	HannXRQ_Chr1 0g0282371	Putative ribosomal protein S11	downstream_gene _variant	6054
seed_hus k_ratio	S10_244 09447	HannXRQ_Chr1 0g0282361	Putative PGG domain	downstream_gene _variant	81280
seed_hus k_ratio	S10_254 68639	HannXRQ_Chr1 0g0282611	Putative protein kinase-like domain Leucine-rich repeat-containing N-terminal	upstream_gene_v ariant	33872

In particular, SNP S14_14459481 associated with the seed area was located 63kb away from gene BSK1 encoding BR-signaling kinase 1, a receptor that is involved in brassinosteroid signaling. Brassinosteroids are the plant hormones that regulate growth and development as well as seed formation (Clouse, 2011). S11_76005476 SNP associated with the seed area located 3kb upstream of the gene encoding SAUR-like auxin-responsive protein family SAURs were reported to act in auxin-responsive cell elongation (Stortenbeker & Bemer, 2019). Gene encoding probable Translation elongation factor EF1B was located 47kb downstream the S06_1852907 associated with seed area. Previously it was demonstrated that EF1B is involved in the control of cell wall biosynthesis in thale cress (Hossain et al., 2012). For the husk area, an SNP S10_222897158 was located close to the gene encoding Putative transcription factor GRAS Scarecrow-like 29. GRAS Transcription Factors were shown to affect cell growth via gibberellin signaling (Cenci & Rouard, 2017). Unfortunately, previous studies aimed at scanning genetic markers associated with seed size-related traits were performed at a low mapping resolution not allowing for to identification of candidate genes. The only exception was the most recent study

(Reinert et al., 2020), which identified 11 genes as strong candidates for trait control. Among them were genes encoding proteins involved in ubiquitination, glycosyltransferases, and sulfate anion transporters.

6.5 Conclusions

This part of the study lists only preliminary results of the association mapping seed-related traits as this project has been initiated, a new genome assembly of sunflower has been released and became publicly available. Also additional phenotype measurements during different years and/or locations are needed to estimate the genotype–environment interactions for such complex traits. Nevertheless, these preliminary results provided some insights on seed-related traits mapping in the sunflower diversity panel from three collections. To perform mapping, first, information on seed-related traits was collected by means of X-ray radiography followed by image analysis. Phenotype data analysis revealed significant differences in trait distribution between the studied collections, which could be explained by the different breeding directions applied to them. Second, genetic diversity analysis and comparison of the genetic diversity with the international lines were performed. It was established that the diversity of cultivated sunflower lines covers the diversity of the cultivated lines from other international collections. Additionally, a significant genetic difference between wild and cultivated sunflowers, independent of the source collections, was identified. Association mapping revealed loci associated with the seed size-related traits. Further, scanning the genetic regions carrying associated markers reveal several potential candidate genes that could be involved in the analyzed trait expression. It should be noted that the identified genetic markers explained not more than seven percent of phenotypic variance. Thus, since the studied traits apparently are under complex polygenic control, the genomic selection approaches could be more applicable to this trait than marker-assisted ones.

Chapter 7. Conclusions and future perspectives

This study presents an effort to identify genetic markers associated with valuable traits in rapeseed and sunflower crops. Further, as this study included, for the first time, full genome genotyping of a substantial number of diverse rapeseed and sunflower lines grown in Russia, I further performed a population structure analysis and diversity comparison of the Russian ~~plan~~ material with those used worldwide.

The main conclusions of the work are as follows: 

- 1) Association mapping of the glucosinolate content in rapeseed revealed 7 SNP markers that explained up to 23% of the phenotypic variance. The identified SNP markers are located close to the new candidate genes that could be potentially involved in the control of glucosinolate composition.
- 2) The QTL mapping of oil quality traits in Russian plant material identified novel SNP markers  or previously reported *Tph1*, *Tph2*, and *Ol* loci associated with tocopherol composition and oleic acid content, respectively. The respective genetic markers were validated using independent plant samples.
- 3) Association mapping of seed, husk and seed to husk area traits conducted based on 601 Russian sunflower lines revealed 25 SNP markers associated with the seed-related traits. A candidate gene analysis revealed new candidate genes involved in the seed-related trait expression.
- 4) Analysis of the population structure of VNIIMK collection of the rapeseed revealed the strong population clustering, which is partially explained by the phenotypical and phenological features. Comparative analysis of the genetic diversity revealed genetic differences potentially underlying the separation between the international and VNIIMK accessions.
- 5) Analysis of the genetic diversity of the sunflower accessions from VNIIMK, VIR and AGROPLASMA collections did not reveal any significant population structure. Further, a joint population structure analysis of a large collection of international and Russian accessions from VNIIMK, VIR and AGROPLASMA

collections identified no significant genetic structure differences between cultivated accession from International and Russian accession. However, we observed significant genetic differences between the wild and cultivated sunflowers independent of the collection origin.

The genetic markers of glucosinolate content could be used to develop approaches for marker-assisted selection for oil and oilcake quality improvement right after the validation of these markers on the independent plant sample of rapeseeds. Genetic markers associated with *Tph1* and *Tph2* were validated by two independent approaches and could be used to develop PCR-based systems for MAS with the aim of tocopherol composition prediction. Markers associated with the *Ol* should be additionally validated on the VNIIMK lines that will be used to obtain varieties with high oleic acid content. Genetic markers associated with the seed-related traits could further provide a better understanding of these traits' genetic control. However, our study suggests that considering the low amount of the seed phenotypic variance explained by the markers, an alternative approach such as genomic selection should be applied to predict and select the proper candidate plants for seed-related trait breeding. Finally, all information on the genetic diversity obtained for rapeseed in sunflower lines could be used as a resource for selection scheme design and genetic diversity analysis by the breeders, as all genotyping data associated with the project is freely available.

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