### Julius-Maximilians Universiät Würzburg

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Quantitative genetics - from genome assemblies to neural network aided omics based prediction of quantitative traits

Author: Supervisor:

Jan Alexander Prof. Arthur KORTE

FREUDENTHAL

for the degree of Ph.D.

in the

Research group for evolutionary genomics

GSLS

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**Declaration of Authorship** 

"Wit beyond measure is man's greatest treasure"

Rowena Rawenclaw

### JULIUS-MAXIMILIANS UNIVERSIÄT WÜRZBURG

## Abstract

Faculty Name GSLS

Ph.D.

# Quantitative genetics - from genome assemblies to neural network aided omics based prediction of quantitative traits

by Jan Alexander FREUDENTHAL

The Thesis Abstract is written here (and usually kept to just this page). The page is kept centered vertically so can expand into the blank space above the title too...

# Acknowledgements

The acknowledgments and the people to thank go here, don't forget to include your project advisor...

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## List of Abbreviations

Adadelta Adaptive delta

Adagrad Adaptive Gradient Algorithm

Adam Adaptive Moment estimation

ANN Artificial Neural Network

**AUC** Area Under the Curve

BLUE Best Linear Unbiased Estimator

BLUP Best Linear Unbiased Predictor

**BP** Base Pair

CPU Core Processing Unit

DH Doubled Haploid

DNA DeoxyriboNucleic Acid

DNA RiboNucleic Acid

**EMMA** Efficient Mixed Model Associations

FCL Fully Connected Layer

GBLUP Genomic Best Linear Unbiased Predictor

GD Gradient Descent

**GP G**enomic **P**rediction

GPU Graphical Processing Unit

**GS** Genomic Selection

GUI Graphical User Interface

**GWAIS** Genome Wide Interaction Association Studies

**GWAS** Genome Wide Association Studies

HDF Hierarchical Data Format

IR Inverted Repeat

LCL Locally Connected Layer

LD Linkage Disequilibrium

LMM Linear Mixed Model

LSC Large Single Copy

MLP Multi Layer Perceptron

ML Machine Learning

MSE Mean Square Error

Nadam Nesterov-accelerated Adaptive Moment Estimation

NAG Nesterov Accelerated Momentum

QTL Quantitative Trait Locus

**ReLU** Rectified Linear Units

**RKHS** Reproducing Kernel Hilbert Spaces

RMSE Root Mean Square Error

RMSProp Root Mean Square Propagation

**ROC** Receiver Operating Characteristics

RSS Residual Sum of Squares

SGD Stochastic Gradient Descent

SNP Single Nucleotide Polymorphism

SSC Small Single Copy

TRN TRaiNing subset

TST TeSTing subset

WGS Whole Genome Sequencing

**XOR** eXclusive **OR** 

MBP Mega Base Pair

SRA Sequence Read Archive

NCBI National Center for Biotechnological Information

**GPL** General Public License

HPC High Performance Computing

GEBC Genomic Estimated Breeding Values

GRM Genomic Relationship Matrix

MCMC Markov Chain Monte Carlo

For/Dedicated to/To my...

# 1 Benchmarking of Chloroplast

## **Genome Assembly tools**

This chapter orientates on *Freudenthal* et al., 2019b only the chapters from the publication which the author majorly contributed are included. If not cited otherwise the plots even though they were published along the aforementioned paper, were designed and generated by the author.

#### 1.1 Introduction

#### 1.1.1 Motivation

Certain organelles like mitochondria and chloroplasts contain their own genetic information from which they are able to synthesize certain proteins independent of the core genome. Evolutionary this developed during endosymbiosis. Its underlying theory seeks to explain how eukaryotic cells formed from prokaryotes *Mereschkowsky*, 1905; *Kutschera* and *Niklas*, 2005. The widely acknowledged theory explains how in the early evolution of eukaryotes, other cells were incorporated in those cells from which today's known organelles descent. In the case of a chloroplast this was most likely a photosynthetic bacteria or similar organism *Archibald*, 2015. Until today the structure of chloroplast genomes resembles more that of a prokaryotic genome than that of its eukaryotic host cells. A typical chloroplast genome consists of circular DNA with a size between 120 kBP to 160 kBP *Palmer*, 1985. The first chloroplast have been sequenced as early as 1986 and were isolated from *Marchantia polymorpha* and *Nico Ohyama* et al., 1986; *Shinozaki* et al., 1986. Complete reviews of

the genome structure of chloroplast genomes were written by Green, 2011; Wicke et al., 2011. Chloroplast genomics is widely applied in evolutionary studies Martin et al., 2002; Xiao-Ming et al., 2017. The genome has been reduced in size over the course of evolution through endosymbiotic gene transfer Martin et al., 2002; Deiner et al., 2017. This is still an ongoing process, which can be observed in vitro Bock, 2017; Fuentes et al., 2014; Stegemann and Bock, 2009. Up to 14 % of the the core genome of Arabidopsis thaliana is made up of genes previously located on the chloroplast (fancy citation), while 100-120 genes remain on the chloroplast Wicke et al., 2011 itself, which by far would not be enough to function independently. Chloroplast genomes are much smaller than core genomes e.g A. thaliana has a core genome size of ca. 125 MBP and are highly conserved with a large gene content, therefore polymorphisms are more likely to cause functional changes in the metabolism. Single chloroplast contain up to hundreds of copies of its genome Kumar, Oldenburg, and Bendich, 2014; Bendich, 1987. Photosynthetic activate cells contain multiple chloroplasts, therefore the copy number of the chloroplast genomes is much higher than the number of core genomes per cell which is most cases is 1.

Structurally chloroplast genomes are made up of 4 distinct regions: two inverted repeats (IR) -  $IR_A$  and  $IR_B$  - ranging from 10 kbp to 76 kbp. They divide the genome into two areas: the large single copy (LSC) and the small single copy (SSC) as shown in 1.1 *Palmer*, 1985. Taking into account that the majority of assembly tools has been designed to assemble linear core genomes, the structure of chloroplast genomes is a major obstacle when wanting to assemble those genomes with modern short read technologies, especially solving and aligning the IR *Wang* et al., 2018.

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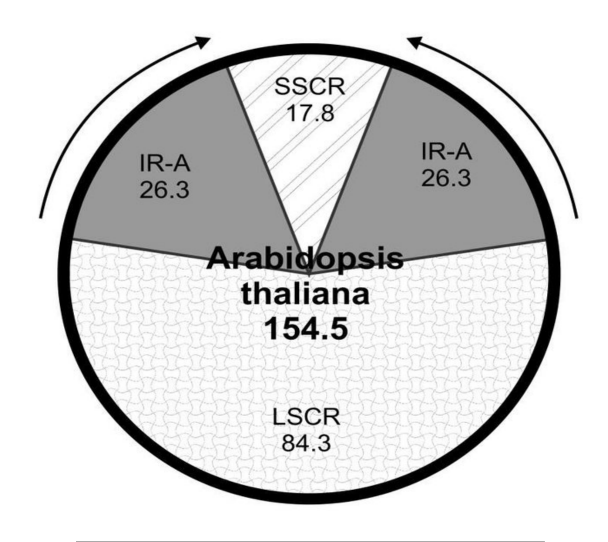


FIGURE 1.1: Structure of the chloroplast genome of *A. thaliana* with SSC-Region and LSC-Region. Length of the genome and its parts in kbp. Graphic from *Olejniczak* et al., 2016

Chloroplast genomes present biologist with many obstacles of which heteroplasmy stands out and immensely complicated genome assemblies and downstream analyses *Corriveau* and *Coleman*, 1988; *Chat* et al., 2002. Heteroplasmy describes the phenomenon of co-existence of multiple versions of the chloroplast genomes in a single organism and single cells of that respective organism. The underlying evolutionary mechanisms behind heteroplasmy are no fully elucidated and eventually existing fitness advantages fueling heteroplasmy cannot be explained satisfactory by evolutionary methods *Scarcelli* et al., 2016. There is a variety of databases containing short read data for species without an chloroplast genome available. For

example NCBI's the short read archive (SRA) *Leinonen* et al., 2010. Because most plant DNA extraction protocols use green leaf tissue as basis, they usually contain a large amount of plastid DNA. Having larger number of assembled and annotated chloroplast genomes publicly available would be beneficial for evolutionary studies and could be a useful addition to bar-coding and super-barcoding *Coissac* et al., 2016 and other biotechnological applications *Daniell* et al., 2016. To achieve this there is a variety of tools available. The study presented in this chapter, assesses the availability, usability and overall performance of 7 of those tools and ultimately makes use of the newly gained insights to attempt *de novo* assemble more than 100 chloroplasts.

# 1.1.2 Extraction of chloroplast reads from whole genome data and general assembly workflow

There is a variety of strategies to assemble chloroplast genomes from raw sequencing data *Twyford* and *Ness*, 2017. In general the process involves three steps: (i) extraction of plastid reads from the WGS data, (ii) assembly of the plastid genome, (iii) solving the circular structure of the chromosome with the IRs. There are two distinct ways to tackle step (i): The first one is to map all the reads to a reference chloroplast *Vinga* et al., 2012, which works reasonably well if there is one available for the same, or at least a closely related species. The second is to make use of the much larger coverage of chloroplast DNA compared to core DNA, with a k-mer analysis *Chan* and *Ragan*, 2013, this is for example done by chloroExtractor*Ankenbrand* et al., 2018. The third way to accomplish this is to combine as done by NOVOPlasty*Dierckxsens*, *Mardulyn*, and *Smits*, 2017. Figure 1.2 shows the general workflow of chloroplast assembly too.

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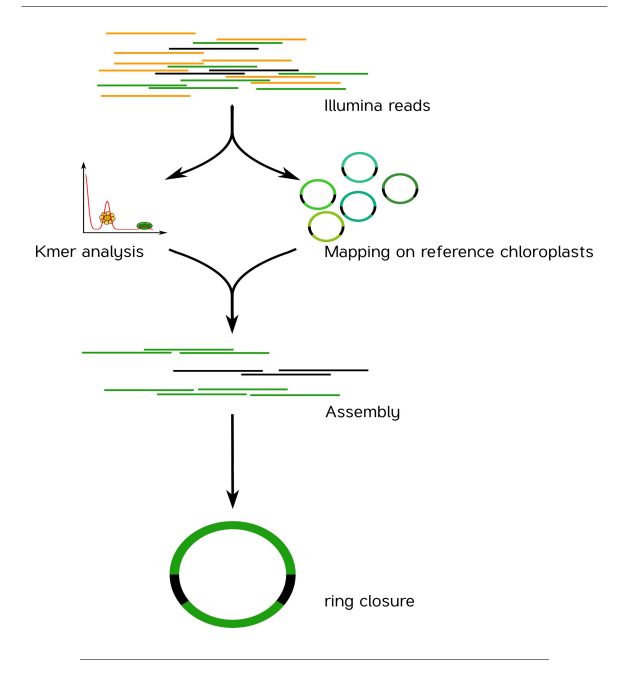


FIGURE 1.2: Standard workflow of chloroplast genome assembly. Graphic from *Ankenbrand* et al., 2018

#### Purpose and scope of benchmarking the landscape of chloroplast assembly tools

The purpose of this study is to provide insights into the landscape of chloroplast assembly tools, to recommend best practices for organelle genome assemblies and to provide *de novo* assemblies for many species and family's without a reference chloroplast available so far. To be included into this study the software including

the source code must be publicly available and published under the terms of a liberal free open source software e.g. GPL or MIT-license. The study was also limited to paired-end Illumina data sets as their sole input source, because they are abundantly available for this benchmark. The seven tools that me the requirements and were therefore included in this study were: chloroExtractor, ORG. Asm, Fast-Plast, IOGA, NOVOPlasty, GetOrganelleand Chloroplast assembly protocol. As later thoroughly described in section 1.3, there are huge differences between those tools and some will not be recommend to be used in scientific applications, while others outstand for most but not all cases.

### 1.2 Material and Methods

#### 1.2.1 Methods

#### Data and code availability

All the source code and data used in the scope of this study is publicly available under the terms of the MIT-License. The source code has been published on github *GitHub Repository for Benchmark Project* and archived on zenodo *Förster* and *Ankenbrand*, 2019. The docker images are available on dockerhub *Docker Hub Group for Benchmark Project* 

#### **Tools**

There were certain requirements to be met to be included into the study as aforementioned. The only technical requirements was, being able to assemble chloroplast genomes from paired-end Illumina reads. The other requirements were due to reproducibility. (i) The software must be open-source and available under the terms of a liberal software license. In the authors opinion obscuring reproducibility

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under paywalls cannot be considered good scientific practice. (ii) It must be a command line tool, GUI-only tools are not suited for highly repetitive automated analyses. In total there were 7 tools that met those requirements: (i) chloroExtractor *Ankenbrand* et al., 2018; (ii) Chloroplast assembly protocol *Sancho* et al., 2018; (iii) GetOrganelle *Jin* et al., 2018; (iv) ORG. Asm *Coissac* et al., 2016; (v) IOGA *Bakker* et al., 2016; (vi) Fast-Plast *McKain* and *Afinit*, 2017; (vii) NOVOPlasty *Dierckxeens*, *Mardulyn*, and *Smits*, 2017. There are other tools available, that did not meet the requirements.

#### Standardization and reproducibility

The main goal of the study was to provided deep insights into the landscape of chloroplast assembly tools. To accomplish that we tried to use the highest standards in bioinformatics when it comes to standardization and reproducibility. Along the study we also wanted to publish easy and ready-to-use versions of all the involved programs, working with standardized input. To accomplish that we decided to make us of containerization in form of docker containers Merkel, 2014 to work with the containers in a closed HPC environments we decided a related software - singularity *Kurtzer*, *Sochat*, and *Bauer*, 2017. Therefore users do not have anything else to do than provide two files one for the forward reads (forward.fq) and one for the reverse reads (reverse.fq). Both files are required to be in FASTQ format. If docker or singularity are installed no further setup is requirement, because all dependencies and packages are installed and run from the containers on any given environment. Besides the individual output files, recording the process of the respective program, all programs write the assembly products in to files called output.fa in FASTA format. For the quantitative and consistency measurements the singularity containers were run on the Julia HPC-Cluster of the University of Würzburg using the SLURM workload manager Jette, Yoo, and Grondona, 2002. All runs for all assemblies were set with a time limit of 48h. This was necessary because some assemblers e.g. IOGA,

if not finishing after at least 12 hours appear to continue in some kind of loop and never finish.

#### 1.2.2 Data

Three different data sets were used for this study. (i) Simulated data from *A. thaliana* chloroplasts. (ii) real data with known reference chloroplast to rate the success of the assemblies. (iii) Novel data sets from NCBI's SRA without a know reference chloroplast to apply the gained knowledge for the *de novo* assembly of more than 100 chloroplasts.

#### Simulated

For first steps in any benchmarking process it is always useful to start with simulated data, where the investigators have full control over all the parameters involved. In the case of the present study the data's input parameters thought to be influential on the outcome where: The read length, the ratio between chloroplast and core genome reads. The simulations were based on data from the TAIR10 genome *Berardini* et al., 2015 and performed using seqkit *Shen* et al., 2016. Ratios simulated were: 0:1, 1:10, 1:1000 and 1:1000, with read length of 150 and 250 bp. The simulated data consisted either of 2 million read pairs or the full data available. The simulation process is documented and the code and the data is available on github and zenodo *Ankenbrand* and *Förster*, 2019

#### Real data set

Real data was selected from the SRA database table 1.1 lists the search terms that had to be met according to the aforementioned search terms.

choice	option	explanation
green plants	Organism	include only photosynthetic plants e.g. no algae
wgs	Strategy	only data from wgs projects included
Illumina	Platform	include only paired-en Illumina reads
biomol DNA	Properties	include only biomol. DNA samples (e.g. no RNA)
paired	Layout	exclude single-end reads
random	Selection	
public	Access	Only publicly available data included

TABLE 1.1: Data selection criteria for real data sets from SRA

In total this resulted in 369 data sets representing a broad variety of the plant kingdom with many different families and genera included.

#### Novel data sets

To assess the performance on assemblies without a published chloroplast on CpBase *CpBase* 105 data sets were selected from SRA, it was emphasized that such data sets were selected, with the next relatives with a reference chloroplast as distantly related as possible in taxonomic terms according to NCBI *NCBI Taxonomy* 

#### 1.2.3 Evaluation

#### Quantitative

Each assembly from each assembler was compared to their respective reference genome by alignment using minimap2 Li, 2018 and based on those alignments scores were calculated following equation 1.1 from 0 to 100 with 100 being a perfect score. Four different metrics contributed equally to the final score. (i) The coverage of the assembled genome compared to the reference genome  $cov_{ref}$  as an estimate for the completeness. (ii) The vice versa case  $cov_{qry}$  as a measure for the correctness of the

assembly. (iii) The success of resolving the IR correct, estimated from the size difference from the reference and the newly assembled genome is:  $min \left\{ \frac{cov_{qry}}{cov_{ref}}, \frac{cov_{ref}}{cov_{qry}} \right\}$ . (iv) The number of total contigs were weighted as  $\frac{1}{n_{contigs}}$  giving a chloroplast with 1 contig the optimal score.

$$score = \frac{1}{4} \cdot \left( cov_{ref} + cov_{qry} + min\left\{ \frac{cov_{qry}}{cov_{ref}}, \frac{cov_{ref}}{cov_{qry}} \right\} + \frac{1}{n_{contigs}} \right) \cdot 100$$
 (1.1)

While it is difficulty to assess the success or failure of assemblies on a continuous scale, equation 1.1 allows for objective and unbiased measurements. SNPs or other small variants do not influence the outcome of the score, because it much more likely that there are due to in-species variation of the plastid's genome and not caused be the assembly, and even if the latter is true it would be difficult to evaluate that.

#### Consistency

In any given bioinformatical application consistency is a desired trait. All software (exceptions excluded) should given repeatedly the same input provide the same output. The same obviously or even more so holds true for assembly tools. To evaluate the reproducibility of the the 7 tools. All the 369 real data sets described in section 1.2.2 has been used twice for assembly with each assembler and scored twice with equation 1.1. The correlation between the first and the second scores and runs was used the measure for the robustness of a program.

1.3. Results

### 1.3 Results

#### 1.3.1 Quantitative

#### Simulated data

The simulated data was assembled and scored with all the tools as described above. Figure 1.3 shows as a tile plot the results for all data sets and assemblers. While at first sight there is no clear correlation with the input data sets. It is obvious that there are grave differences between the assemblers. Two programs namely Chloroplast assembly protocol and IOGA failed to correctly assemble a single chloroplast genome. IOGAeven fails to provide output at all for the majority of the data sets. While those two stand out as a negative example Fast-Plastand GetOrganellestand out as a positive example perfectly or nearly perfectly assembling all the data sets, with GetOrganellesurpassing the performance of Fast-Plast. In the middle of the filed are chloroExtractor, ORG. Asmand NOVOPlastyperforming reasonably well, but sometimes lacking to solve the IRs and the circular structure.

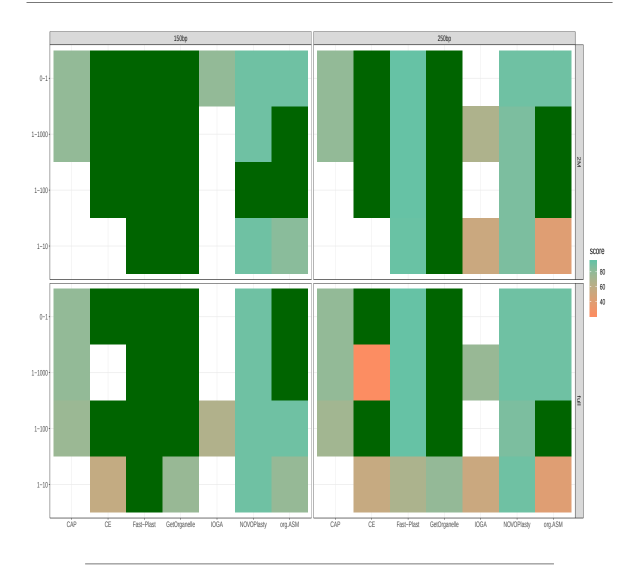


FIGURE 1.3: Results of assemblies executed with simulated data sets.

While there is a significant difference between the assemblers the same is not necessarily true in general for the other varying parameters. While Fast-Plastdeals with the shorter reads of 150 bp much better than with the longer reads of 250 bp. The outcome of the other assemblies does not seem to be influenced by that. There is no difference between the full and the subsampled data sets. And while all assemblers appear to be more challenge by low chloroplast to core genome ratios of 1:10 larger rations do not affect the quality of the result as expected. Table 1.2 shows all the individual results for all data sets and assemblers. For the fields with no entry the respective assembler failed to provide an output.

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TABLE 1.2: Scores of assemblies of simulated data

	data set	CAP	CE	Fast-Plast	GetOrganelle	IOGA	NOVOPlasty	org.ASM
1	sim_150bp.0-1	79.10	100.00	99.48	100.00		91.52	100.00
2	sim_150bp.0-1.2M	79.10	100.00	99.72	100.00	79.10	91.52	91.50
3	sim_150bp.1-10		56.44	100.00	76.98		91.52	78.00
4	sim_150bp.1-10.2M			99.97	100.00		91.52	82.72
5	sim_150bp.1-100	75.72	100.00	99.48	100.00	66.09	91.52	91.50
6	sim_150bp.1-100.2M		100.00	99.47	100.00		100.00	100.00
7	sim_150bp.1-1000	79.10		99.72	100.00		91.52	100.00
8	sim_150bp.1-1000.2M	79.10	100.00	99.72	100.00		91.52	100.00
9	sim_250bp.0-1	79.10	100.00	93.82	100.00		91.52	91.50
10	sim_250bp.0-1.2M	79.10	100.00	93.83	100.00		91.52	91.50
11	sim_250bp.1-10		54.98	68.45	78.89	52.71	91.52	40.20
12	sim_250bp.1-10.2M			93.00	100.00	52.67	87.40	40.20
13	sim_250bp.1-100	72.81	100.00	93.82	100.00		87.40	100.00
14	sim_250bp.1-100.2M		100.00	93.83	100.00		87.40	100.00
15	sim_250bp.1-1000	79.10	21.30	93.83	100.00	76.96	91.52	91.50
_16	sim_250bp.1-1000.2M	79.10	100.00	93.83	100.00	67.55	87.40	100.00

#### Real data sets

Table 1.3 gives the results from the assemblies of 369 data sets with 7 assemblers. Similar to the results of the previous section there is a significant difference between the tools. Likewise GetOrganelleis the most successful assembler by a large margin with 210 of 369 perfectly assembled chloroplast genomes, failing to provide output for only 9 data sets, resulting in a media score > 99. Contrary to GetOrganelle, Chloroplast assembly protocoland IOGAfail to completely assemble a single genome. The performance of Fast-Plastcould be considered reasonably well, completing approximately half as many genomes as GetOrganelleand being the only other tool whose average score is larger than 90. Similar to the trials with the simulated data in chapter 1.3.1 in the middle of the field are chloroExtractor, NOVOPlasty and ORG. Asm.

	assembler	Median	IQR	N_perfect	N_tot
1	CAP	45.25	50.19	0	369
2	CE	56.55	71.50	14	369
3	Fast-Plast	92.80	23.59	113	369
4	GetOrganelle	99.83	20.94	210	360
5	IOGA	71.10	11.21	0	338
6	NOVOPlasty	75.95	48.69	58	369
7	org.ASM	67.35	91.69	46	348

TABLE 1.3: Mean scores of chloroplast genome assemblers

Figure 1.4 emphasizes the large differences between the assemblers shown in table 1.3. The swarm plots show some distinct bands for some assemblers e.g. NOVOPlastyand ORG. Asm, suggesting that multiple assemblies fail to be solved into a single contig genome at a certain point. As thoroughly discussed in section 1.4 just from this swarm plot it is debatable if all the tools could be recommended for the purpose they were designed for.

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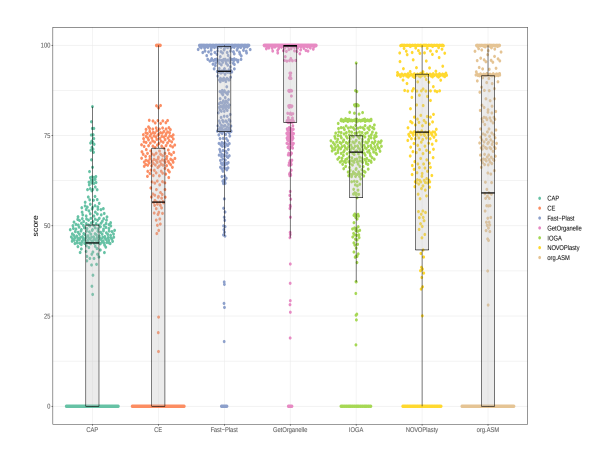


FIGURE 1.4: Box and swarm plots depict the results from the scoring shown in 1.1

#### Consistency

Consistency testing was done by re-running every assembly for the real data sets twice and comparing the scores. chloroExtractorwas the only tool that was 100 % consistent over 2 runs. Followed by Fast-Plastand NOVOPlasty. The consistency plot figure 1.5 for both of them results in an arrowhead shaped plot. With the main differences between the first and second run in the best scores. All other assemblers appear to produce the same output in the two runs except if the fail to complete the assembly at all. This is less pronounced for Chloroplast assembly protocoland GetOrganelleand is a grave issue for ORG. Asmand IOGA.

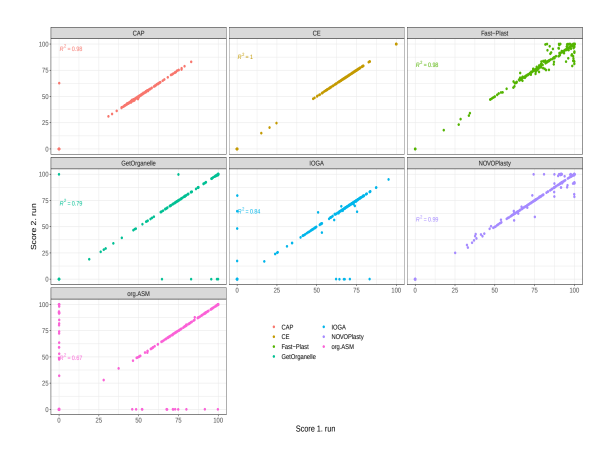


FIGURE 1.5: Swarm plots depict the results from the scoring shown in 1.1 for two independent runs for each assembler on each of the data sets

#### **Novel**

The final assessment in the scope of the study was to test the assemblers on novel data sets without a published chloroplast. This step is important for two reasons. (i) There could be the possibility that certain tools perform well on known chloroplasts because they somehow of knowledge over those and simply apply it during those assemblies and therefore the previous results will not be able to generalize on unknown data. (ii) To apply and test the gained insights in the hope of providing the scientific community with a larger variety of published chloroplast genomes. Again the most successful assembler was GetOrganelle, with 49 out of 105 data sets completely assembled. Lacking a reference genome the success had to be defined differently and equation 1.1 was not suitable to evaluate the novel assemblies.

Metrics influencing the score of the novel assemblies were the number of contigs, solving the IRs and the size of the SSC and LSC. This known to the authors might be very biased and not true for all chloroplast and assumes that all chloroplast genomes follow the general structure for their genomes described in chapter 1.1. Figure 1.6 compares the results for the assemblies with at least one successful assembly.

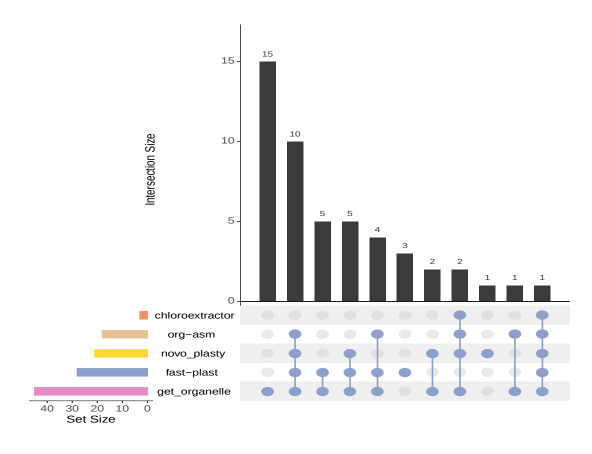


FIGURE 1.6: Upset plot comparing the success rates of the different assemblers

# 1.4 Discussion

The study compromised of two goals. (i) to assess the overall performance of a variety of tools designed specifically for the assembly of circular chloroplast genomes from paired-end Illumina data and (ii) to *de novo* assemble a variety of yet unpublished chloroplast genomes from existing data. To accomplish the first goal 16 simulated and 369 simulated data sets were used adding up to a total of 5166 assemblies

for the real data sets and 112 for the simulated data, along 735 assemblies for the novel data sets, thus underlying the statistical powers of this benchmarking study. The most successful tools were GetOrganelle and Fast-Plast which are recommended to be used complementary, because as shown in figure 1.7 they succeed for the most data sets compared to other assemblers and accomplish to satisfactory assemble chloroplast genomes where the other fails. If both of them fail it might be worthwhile to repeat the runs because other results could be expected as shown in the scatter plots of figure 1.5, especially Fast-Plastmight be able to improve the previously achieved sore. Only if both of them fail it might, even though improbable, that NOVOPlastymight lead to a successful assembly. The other assemblers should be used with caution. While chloroExtractormight be good for a quick overview due to its relatively low demand in computational time *Freudenthal* et al., 2019b; Chloroplast assembly protocol, ORG. Asmand IOGAare not be recommended to used as the primary tools in organelle genome assembly projects, as used in this study.

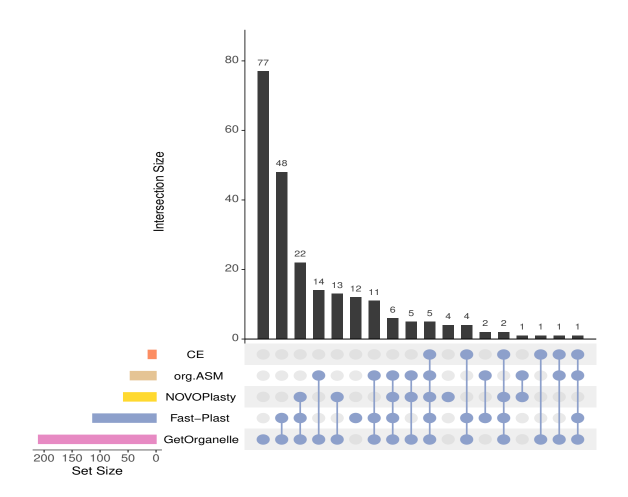


FIGURE 1.7: Upset plot showing the intersections of success rates between assemblers. A successful assembly was defined with a score > 99 according to equation 1.1. The colored, horizontal barplot indicates the total number of successful assemblies for an individual tool. The black, vertical barplot gives the magnitude of the intersection between the assemblers indicated with the dot. Therefore there first bar is to be interpreted as follows: 77 data sets were only successfully assembled by chloroExtractor, likewise 48 genomes were assembled completely by GetOrganelle and Fast-Plast and so on.

It might be possible that overall performance of a specify tool might change significantly by fine tuning the input parameters of the tool, which was purposely not done in the scope of the present study, because this study was designed to mimic

the behavior have end-users and not developers of such tools and the assumption is proposed that end-users with little experience in bioinformatics would be inclined to use the basic configurations of such a tool.

While there is a huge differences for all assemblers they are presented with the same challenges and the bottlenecks are similar for each of them, the success rate of passing those differs however. Figure 1.8 shows the alignment of the 7 chloroplast genomes of Oryza brachyantha a grass distantly related to cultivated rice Orayza sativa and the respective reference genome. For the need of a linear representation of the circular genome the convention is to present chloroplast genomes in the order LSC - IRa - SSC -IRb. O. brachyantha was chosen because multiple tools successfully or at least almost assembled the full genome. Only Chloroplast assembly protocolis singled out, which only managed to assemble a few fragments on the SCC and the IRs on many contigs. A common mistake is to return 3 contigs as IOGA did. They represent the LSC the SSC and one IR but failed to resolve those regions into a one circular contig. GetOrganelle and Fast-Plast were able to reproduce the structure of the reference, while chloroExtractor flipped the LSC and NOVOPlastyand ORG. Asm were not able to construct the single contig into the conventional structure. All of this are common mistakes appearing more or less rare in all the assemblers. This could be a good starting point for the developers to further improve their tools. In this example all but Chloroplast assembly protocol were able to construct all the parts of the chloroplast's genome, and the main mistake was to resolve the structure of the genome into a circular, one contig version.

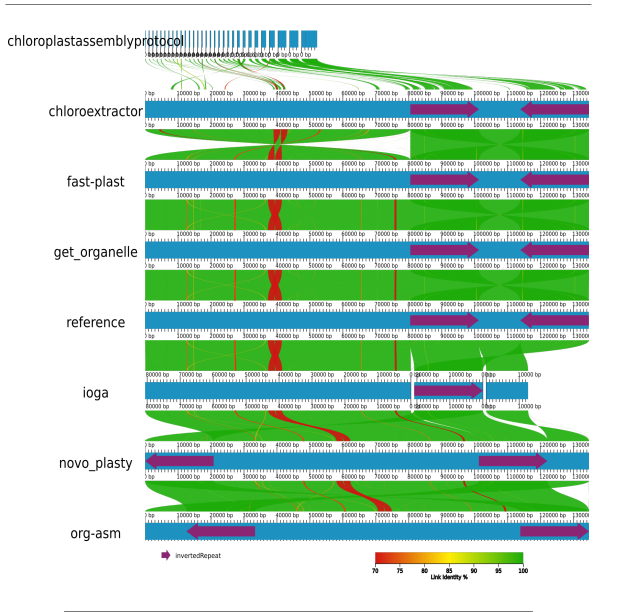


FIGURE 1.8: AliTV plot *Ankenbrand* et al., 2017 from *Freudenthal* et al., 2019b showing the alignments of *Oryza brachyantha* chloroplast genomes fro all 7 assemblers. Regions in adjacent assemblies are connected with colored ribbons for similar regions in alignment with the identity coded as described in the legend. The purple arrows indicate the IR regions

# 1.5 Conclusion & outlook

Organelle genomics is promising field in plant genetics. As described in section 1.1 chloroplast genomes are well-suited for applications in evolutionary sciences, taxonomy and barcoding applications. Alike its mother branch genomics for comparative chloroplast genomics its just as crucial to obtain high quality genomes. And the quality is mainly influenced by two major factors: the quality of the genome sequencing protocol and the quality of the assembly. As shown the latter varies massively between tools and not all tools are recommend equally from the conclusions drawn for the above experiments. All tools have room for improvement, this is meant to criticize the respectable work of the developers, but to encourage them to further develop tools and publish them under terms of liberal software licenses for the greater benefit of the entire scientific community.

# 2 Understanding the haplotype structure of Arabidopisis thaliana

## 2.1 Introduction

Recombination and LD in *A. thaliana Kim* et al., 2007 LD in *A. thaliana Nordborg* et al., 2002 Evolution of selfing *Tang* et al., 2007 Evolution and genetic differentiation among relatives of Arabidopsis thaliana *Koch* and *Matschinger*, 2007 FLC haplotypes *Li* et al., 2014

- 2.2 Haplotyping of A. thaliana
- 2.3 Results
- 2.4 Disucssion

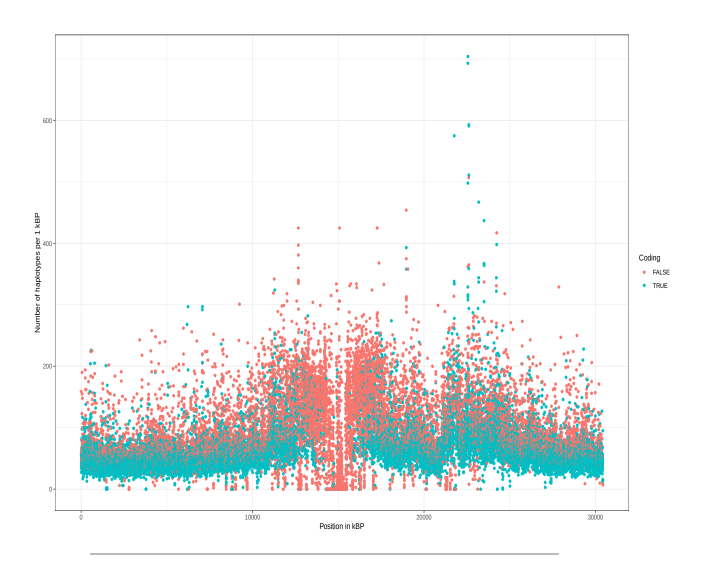


FIGURE 2.1: The number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

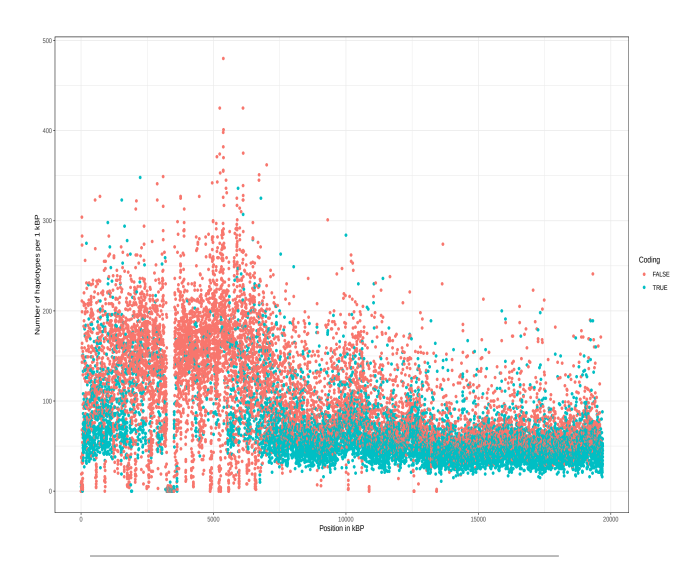


FIGURE 2.2: Number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

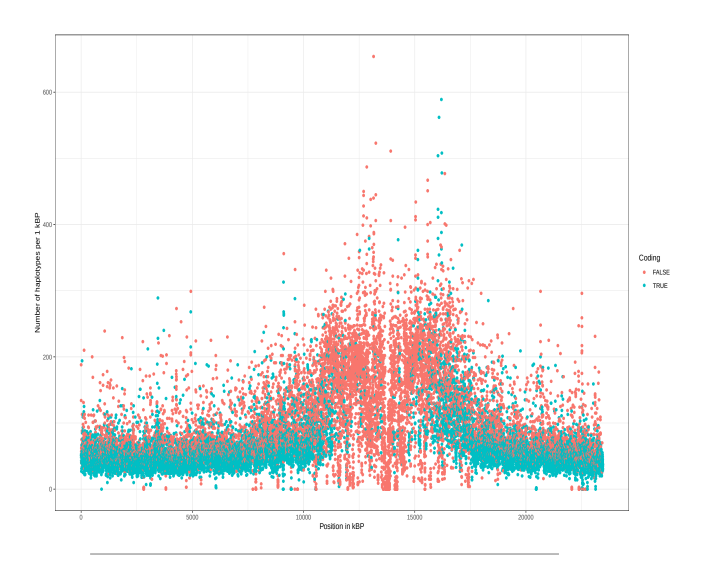


FIGURE 2.3: Number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

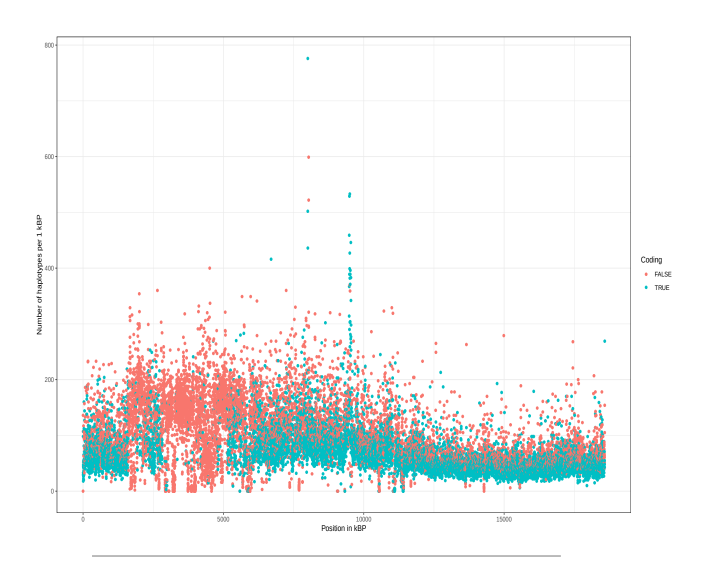


FIGURE 2.4: Number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

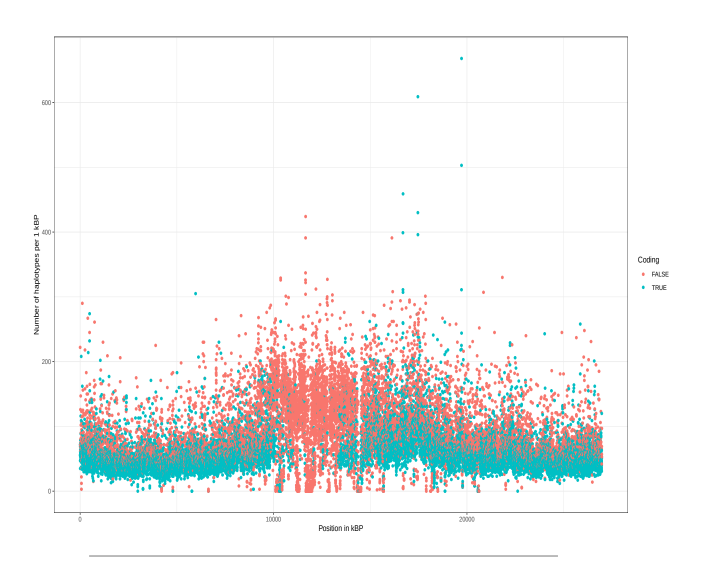


FIGURE 2.5: Number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

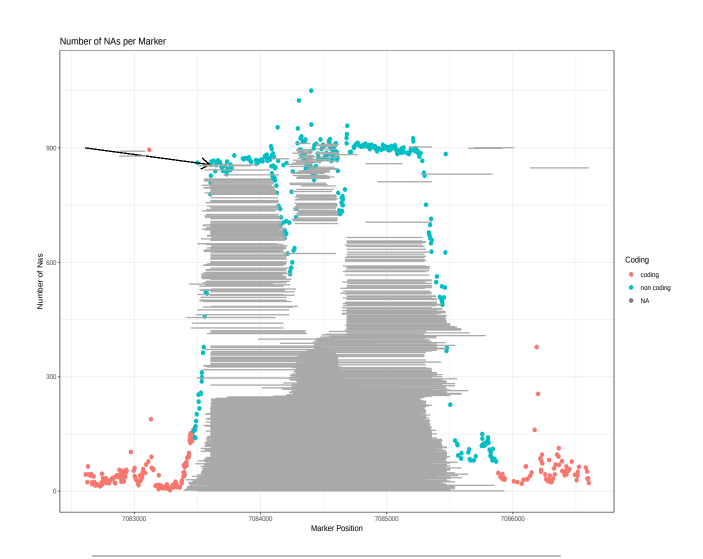


FIGURE 2.6: Number of segregating haplotypes with a polymorphism in at least one position over a stretch of 1 kBP.

# 3 GWAS-Flow a gpu-accelerated software for large-scale genome-wide association studies

The following chapter has been published in a similar version on the bio $\chi$ iv preprint server *Freudenthal* et al., 2019a and has been submitted for publication to Oxford Bioinformatics. The experiments and the software have been designed and conducted author. The manuscript has been prepared by the author, with minor corrections from Prof. Arthur Korte & Dominik Grimm. All authors approved of the final manuscript.

### 3.1 Introduction

Genome-wide association studies, pioneered in human genetics *Hirschhorn* and *Daly*, 2005 in the last decade, have become the predominant method to detect associations between phenotypes and the genetic variations present in a population. Understanding the genetic architecture of traits and mapping the underlying genomic polymorphisms is of paramount importance for successful breeding both in plants and animals, as well as for studying the genetic risk factors of diseases. Over the last decades, the cost for genotyping have been reduced dramatically. Early GWAS consisted of a few hundred individuals which have been phenotyped and genotyped on a couple of hundreds to thousands of genomic markers. Nowadays, marker density

for many species easily exceed millions of genomic polymorphisms. Albeit commonly SNPs are used for association studies, standard GWAS models are flexible to handle different genomic features as input. The Arabidopsis 1001 genomes project features for example 1135 sequenced Arabidopsis thaliana accessions with over 10 million genomic markers that segregate in the population *Alonso-Blanco* et al., 2016. Other genome projects also yielded large amounts of genomic data for a substantial amount of individuals, as exemplified in the 1000 genomes project for humans Siva, 2008, the 2000 yeast genomes project or the 3000 rice genomes project Li, Wang, and Zeigler, 2014. Thus, there is an increasing demand for GWAS models that can analyze these data in a reasonable time frame. One critical step of GWAS is to determine the threshold at which an association is termed significant. Classically the conservative Bonferroni threshold is used, which accounts for the number of statistical tests that are performed, while many recent studies try to significance thresholds that are based on the false-discovery rate (FDR) Storey and Tibshirani, 2003. An alternative approach are permutation-based thresholds Che et al., 2014. Permutation-based thresholds estimate the significance by shuffling phenotypes and genotypes before each GWAS run, thus any signal left in the data should not have a genetic cause, but might represent model mis-specifications or uneven phenotypic distributions. Typically this process is repeated hundreds to thousands of times and will lead to a distinct threshold for each phenotype analyzed Togninalli et al., 2017. The computational demand of permutation-based thresholds is immense, as per analysis not one, but at least hundreds of GWAS need to be performed. Here the main limitation is the pure computational demand. Thus, faster GWAS models could easily make the estimation of permutation-based thresholds the default choice.

3.2. Methods

### 3.2 Methods

#### **GWAS Model**

The GWAS model used for GWAS-Flow is based on a fast approximation of the linear-mixed-model described in Kang et al., 2010; Zhang et al., 2010, which estimates the variance components  $\sigma_g$  and  $\sigma_e$  only once in a null model that includes the genetic relationship matrix, but no distinct genetic markers. These components are thereafter used for the tests of each specific marker. Here, the underlying assumption is, that the ratio of these components stays constant, even if distinct genetic markers are included into the GWAS model. This holds true for nearly all markers and only markers which posses a big effect will alter this ratio slightly, where now  $\sigma_g$  would become smaller compared to the null model. Thus, the p-values calculated by the approximation might be a little higher (less significant) for strongly associated markers.

#### The GWAS-Flow Software

The GWAS-Flow software was designed to provide a fast and robust GWAS implementation that can easily handle large data and allows to perform permutations in a reasonable time frame. Traditional GWAS implementations that are implemented using Python *Van Rossum* and *Drake Jr*, 1995 or R R Core Team, 2019 cannot always meet these demands. We tried to overcome those limitations by using TensorFlow Abadi et al., 2015, a multi-language machine learning framework published and developed by Google. GWAS calculations are composed of a series of matrix computations that can be highly parallelized, and easily integrated into the architecture provided by TensorFlow. Our implementation allows both, the classical parallelization of code on multiple processors (CPUs) and the use of graphical processing units (GPUs). GWAS-Flow is written using the Python TensorFlow API. Data import is done with pandas McKinney, 2010 and/or HDF5 for Python Collette, 2013. Preprocessing of the data (e.g filtering by minor Allele count (MAC)) is performed with numpy

Oliphant, 2006. Variance components for residual and genomic effects are estimated with a slightly altered function based on the Python package *limix Lippert* et al., 2014. The GWAS model is based on the following linear mixed model that takes into account the effect of every marker with respect to the kinship:

$$Y = \beta_0 + X_i \beta_i + u + \epsilon, u \sim N(0, \sigma_g K), \epsilon \sim N(0, \sigma_e I)$$
(3.1)

From this LMM the residual sum of squares for marker i are calcucated as descirebed in 3.2

$$RSS_i = \sum Y - (X_i \beta_0 + I_i \beta_1) \tag{3.2}$$

The residuals are used to calculate a p-value for each marker according to an overall F-test that compares the model including a distinct genetic effect to a model without this genetic effect:

$$F = \frac{RSS_{env} - R1_{full}}{\frac{R1_{full}}{n-3}} \tag{3.3}$$

Apart from the p-values that derive from the F-distribution, GWAS-Flow also report summary statistics, such as the estimated effect size ( $\beta_i$ ) and its standard error for each marker.

#### Calculation of permutation-based thresholds for GWAS

To calculate a permuation-based threshold, we essentially perform n repetitions (n > 100) of the GWAS on the same data with the sole difference that before each GWAS we randomize the phenotypic values. Thus any correlation between the phenotype and the genotype will be broken and indeed for over 90% of these analyses the estimated pseudo-heritability is close to zero. On the other hand, the phenotypic distribution will stay unaltered by this randomization. Hence, any remaining signal in the GWAS has to be of a non-genetic origin and could be caused by e.g. model mis-specifications. Now we take the lowest p-value (after filtering for the

3.2. Methods 35

desired minor allele count) for each permutation and take the 5% lowest value as the permutation-based threshold for the GWAS.

#### Benchmarking

For benchmarking of GWAS-Flow we used data from the *Arabidopsis* 1001 Genomes Project *Alonso-Blanco* et al., 2016. The genomic data we used were subsets between 10,000 and 100,000 markers. We chose not to include subsets that exceed 100,000 markers, because there is a linear relationship between the number of markers and the computational time demanded, as all markers are tested independently. We used phenotypic data for flowering time at ten degrees (FT10) for A. thaliana, published and downloaded from the AraPheno database Seren et al., 2016. We downand up-sampled sets to generate phenotypes for sets between 100 and 5000 accessions. For each set of phenotypes and markers we ran 10 permutations to assess the computational time needed. All analyses have been performed with a custom R script that has been used previously *Togninalli* et al., 2017, GWAS-Flow using either a CPU or a GPU architecture and GEMMA Zhou and Stephens, 2012. GEMMA is a fast and efficient implementation of the mixed model that is broadly used to perform GWAS. All calculations were run on the same machine using 16 i9 virtual CPUs. The GPU version ran on an NVIDIA Tesla P100 graphic card. Additionally to the analyses of the simulated data, we compared the times required by GEMMA and both GWAS-Flow implementations for > 200 different real datasets from A. thaliana that have been downloaded from the AraPheno Seren et al., 2016 database and have been analyzed with the available fully imputed genomic dataset of ca. 10 million markers, filtered for a minor allele count greater five.

# 3.3 Results

The two main factors influencing the computational time for GWAS are the number of markers incorporated in such an analysis and the number of different accessions, while the latter has an approximate quadratic effect in classical GWAS implementations Zhou and Stephens, 2012. Figure 1A shows the time demand as a function of the number of accessions used in the analysis with 10,000 markers. The quadratic increase in time demand is clearly visible for the custom R implementation, as well as for the CPU-based GWAS-Flow implementation and GEMMA. The GWAS-Flow implementation and GEMMA clearly outperforms the R implementation in general, while for a small number of accessions GWAS-Flow is slightly faster then GEMMA. For the GPU-based implementation the increase in run-time with larger sample sizes is much less pronounced. While for small (< 1,000 individuals) data, there is no benefit compared to running GWAS-Flow on CPUs or running GEMMA, the GPU-version clearly outperforms the other implementations if the number of accessions increases. Figure 1B shows the computational time in relation to the number of markers and a fixed amount of 2000 accessions for the two different GWAS-Flow implementations. Here, a linear relationship is visible in both cases. To show the performance of GWAS-Flow not only for simulated data, we also run both implementations on more than 200 different real datasets downloaded from the Ara-Pheno database. Figure 1C shows the computational time demands for all analyses comparing both GWAS-Flow implementation to GEMMA. Here, the CPU-based GWAS-Flow performs comparable to GEMMA, while the GPU-based implementation outperforms both, if the number of accessions is above 500. Importantly all obtained GWAS results (p-values, beta estimates and standard errors of the beta estimates) are nearly (apart from some mathematical inaccuracies) identical between the three different implementations.

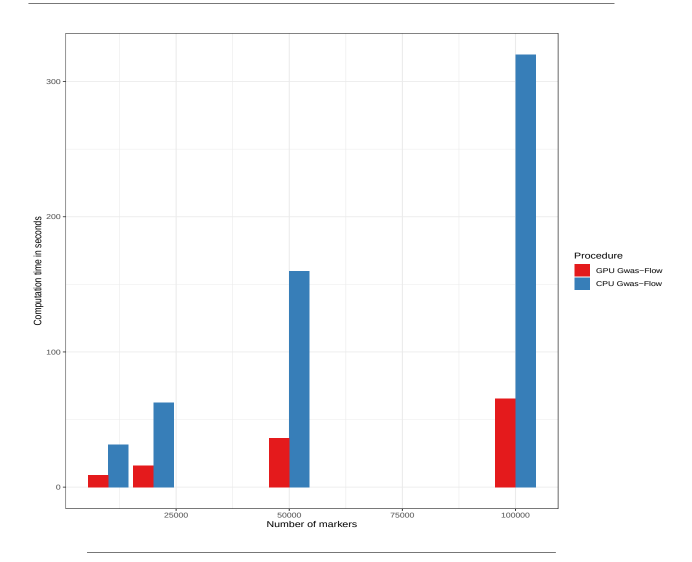


FIGURE 3.1: Computational time as a function of the number of genetic markers with constantly 2000 accessions for both GWAS-Flow versions

# 3.4 Disucssion

We made use of recent developments of computational architecture and software to cope with the increasing computational demand in analyzing large GWAS datasets. With GWAS-Flow we implemented both, a CPU- and a GPU-based version of the classical linear mixed model commonly used for GWAS. Both implementations outperform custom R scripts on simulated and real data. While the CPU-based version performs nearly identical compared to *GEMMA*, a commonly used GWAS implementation, the GPU-based implementation outperforms both, if the number of individuals, which have been phenotyped, increases. For analyzing big data, here the

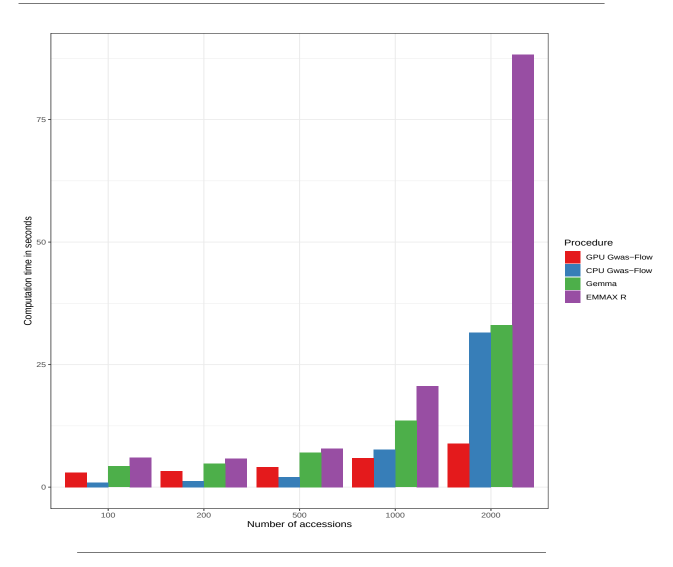


FIGURE 3.2: Computational time as a function of the number of accessions with 10000 markers each.

main limitation would be the RAM of the GPU, but as the individual test for each marker are independent, this can be easily overcome programmatically. The presented GWAS-Flow implementations are markedly faster compared to custom GWAS scripts and even outperform efficient fast implementations like *GEMMA* in terms of speed. This readily enables the use of permutation-based thresholds, as with GWAS-Flow hundred permutations can be performed in a reasonable time even for big data. Thus, it is possible for each analyzed phenotype to create a specific, permutation-based threshold that might present a more realistic scenario. Importantly the permutation-based threshold can be easily adjusted to different minor allele counts, generating different significance thresholds depending on the allele

count. This could help to distinguish false and true associations even for rare alleles. GWAS-Flow is a versatile and fast software package. Currently GWAS-Flow is and will remain under active development to make the software more versatile. This will e.g. include the compatibility with TensorFlow v2.0.0 and enable data input formats, such as PLINK *Purcell* et al., 2007. The whole framework is flexible, so it is easy to include predefined co-factors e.g to enable multi-locus models *Segura* et al., 2012 or account for multi-variate models like the multi-trait mixed model *Korte* et al., 2012. Standard GWAS are good in detecting additive effects with comparably large effect sizes, but lack the ability to detect epistatic interactions and their influence on complex traits *Mckinney* and *Pajewski*, 2012; *Korte* and *Farlow*, 2013. To catch the effects of these gene-by-gene or SNP-by-SNP interactions, a variety of genome-wide association interaction studies (GWAIS) have been developed, thoroughly reviewed in *Ritchie* and *Van Steen*, 2018. Here, GWAS-Flow might provide a tool that enables to test the full pairwise interaction matrix of all SNPs. Although this might be a statistic nightmare, it now would be computationally feasible.

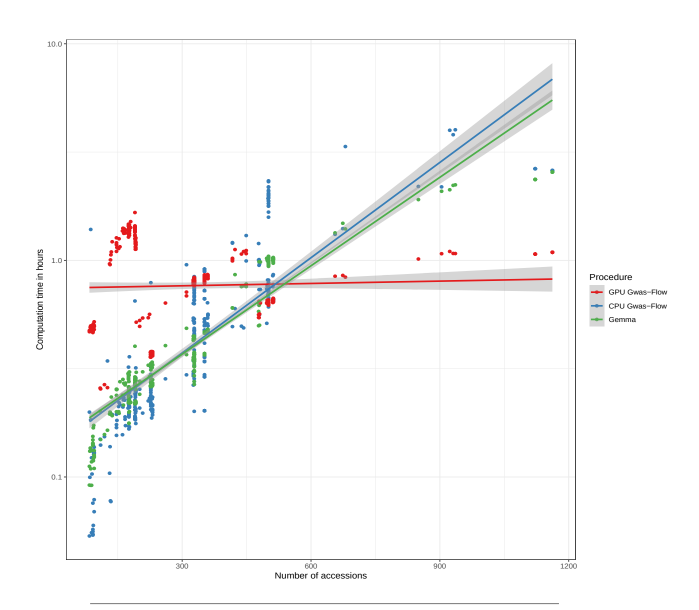


FIGURE 3.3: Comparison of the computational time for the analyses of > 200 phenotypes from *Arabidopsis thaliana* as a function of the number of accessions for *GEMMA* and the CPU- and GPU-based version of GWAS-Flow. GWAS was performed with a fully imputed genotype matrix containing 10.7 M markers and a minor allele filter of MAC > 5

# 4 Genomic prediction of phenotypic values of quantitative traits using artificial neural networks

#### 4.1 Introduction

# 4.1.1 A brief history of machine learning

#### Basic perceptron model

While machine learning, neural networks, deep learning became essential tools for many applications in more recent years, their mathematical principals date back to the early 1950s and 1960s. Figure 4.1 schematically show the basic perceptron model as proposed by Rosenblatt, which was designed to mimic the information flow in biological nervous systems *Rosenblatt*, 1961

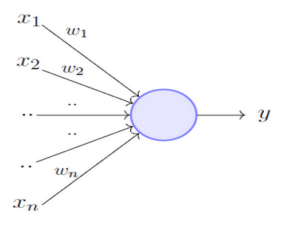


FIGURE 4.1: Basic perceptron model as proposed by Rosenblatt

This basic perceptron, which contrary to perceptrons used nowadays does not have an activation function, takes n binary inputs  $x_1, x_2, ..., x_n$  and produces a single, likewise binary, output y after being processed by the perceptron or neuron. To achieve this Rosenblatt introduced the concept of weights which indicated a certain relative importance to the outcome of the output.  $w_1, w_2...w_n$ . The output y is determined by the weighted sum of the weights and biases  $\sum_i w_i x_i$ . If a certain threshold value is met the neuron is either activated and outputs 1 or not and outputs 0. This is algebraically represented in 4.1

$$0 = \text{if } \sum_{i}^{n} w_{j} x_{i} - \theta \le 0$$

$$1 = \text{if } \sum_{i}^{n} w_{i} x_{i} - \theta > 0$$

$$(4.1a)$$

$$1 = \text{if } \sum_{i=1}^{n} w_{i} x_{i} - \theta > 0$$
 (4.1b)

Next to the weights  $w_n$  and the inputs  $x_n$  a third term  $\theta$  is introduced in equation 4.1 which represents the activation threshold in per definition is negative. A single perceptron is a linear classifier and can only be trained on linearly separable functions and can used as shown by Rosenblatt, 1961 to solve simple logical operations as AND, OR and not. The simple perceptron fails, due to non-linearity, to perform XOR operations as shown by Marvin and Seymour, 1969. This discovery let to a near still stance in the research of artificial neural networks in the 1970s. This time period as now often referred to as the first AI-winter. Another reason that massively hindered the applications and research of machine learning during that time, was the compared to modern times incredibly small amount of computational power available *Nguyen* and *Widrow*, 1990.

More complex decision making, like solving XOR problems, requires more complex structures than a single perceptron. Continuing the trend of mimicking human neural networks, multiple artificial neurons are stacked into layers and these layers, are connected to each other allowing communication between the many perceptrons

4.1. Introduction 43

in a such generated network. Figure 4.2 shows schematically the basic structure of such a network, now container three types of layers. (i) the input layer, (ii) one or more hidden layers and (iii) one output layer, which in this case only consists of one neuron.

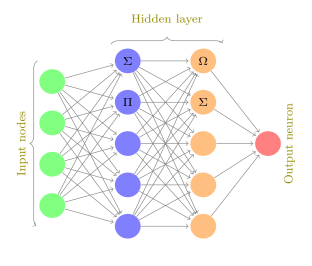


FIGURE 4.2: Schematic layout of a simple multi-layer perceptron

In the sample layout of figure 4.2 the neurons in the first column weigh the inputs and pass those the neurons on the second layer. In this case all neurons on the first layer or connected to all neurons on the second layer, such layers are referred to fully-connected layers (FLC), and their resulting networks are often called multilayer perceptrons (MLP). This architecture enables the network to perform more complex calculations and result in more abstract decisions than single neurons or single layer architectures.

#### **Activation functions**

The neurons discussed so far are only capable of outputting binary results. Either 0 or 1, depending on the threshold values being met or not. For more complex estimations it is desirable that small changes in the input also result in small changes of the output. This requirement can not be met with binary outputs. Activation functions for a given node provides rules for the output in accordance to the inputs *Žilinskas*, 2006.

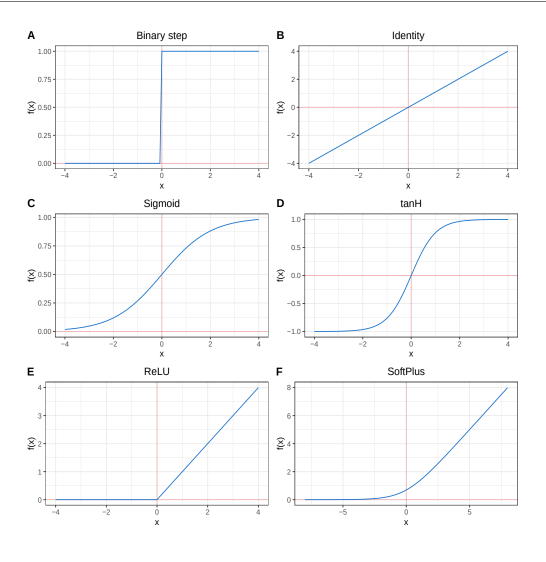


FIGURE 4.3: Popular activation functions used in neural networks. A Binary step activation function. **B** Identity activation function. **C** Sigmoid or logistic activation function. **D** tangens hyperbolicus activation function. **E** rectified linear units activation function . **F** SoftPlus activation function.

Figure 4.3 **A** shows six of the most commonly used activation functions *Warner* and *Misra*, 1996. The simplest one was introduced, is the binary step activation function equation 4.2, which properties have been discussed along the perceptron model. All other activation produce continuous outputs from any given input. Basically any mathematical function can serve as an activation function in neural nets, starting with a simple identity function 4.3, 4.3 **B**. Sigmoid figure 4.3 **C**, equation 4.4 and tanh figure 4.3 **D**, equation 4.5, when  $x \to \infty$  or  $x \to -\infty$  they have similar properties to the binary function, but produce continuous output around 0.

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$$f(x) = \sigma(x) = \begin{cases} 0 \text{ for } x < 0\\ 1 \text{ for } x \ge 0 \end{cases}$$

$$(4.2)$$

$$f(x) = \sigma(x) = x \tag{4.3}$$

$$f(x) = \sigma(x) = \frac{1}{1 + e^{-x}} \tag{4.4}$$

$$f(x) = \sigma(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}}$$
 (4.5)

$$f(x) = \sigma(x) = \begin{cases} 0 \text{ for } x < 0\\ x \text{ for } x \ge 0 \end{cases}$$
 (4.6)

$$f(x) = ln(1 + e^x) \tag{4.7}$$

ReLU (equation 4.6) and the softplus (equation 4.7) share similar properties as well, the latter one being a smoothed version of ReLU. Rectifiers as activation functions have been introduced in 2000s *Hahnloser* et al., 2000 and have since then overtaken all others as the most popular activations functions in neural networks and deep learning today *LeCun*, *Bengio*, and *Hinton*, 2015 and they have proven to be superior in deep-learning algorithms that sigmoid or logistic functions. One of the advantages leading to the superiority of ReLUs is that with randomly initialized weights only half of the ReLU neurons are activated, compared to tanh and sigmoid activation *Glorot*, *Bordes*, and *Bengio*, 2011. All activation functions shown in figure 4.3, but the binary step function, share one common property: a small change of the input weight will result in small changes in the output, while a small change of the input for the binary step function leads to either no or a complete change of the output. This property is, as described below, is an important prerequisite for networks being able to learn.

#### Gradient descent algorithm

Let the network shown in 4.2 be for the classification of a arbitrary phenotype like blue petals with  $x_1...x_4$  on the input layers being genetic markers as features. And the output layer displaying a value from 0 to 1, meaning yes: blue petals from 0 - 0.5 and no blue petals from 0.5 to 1. To quantify how well the network performs on achieving that goal a loss function is applied *Schmidhuber*, 2015. There is a large variety of different loss functions available for neural networks like mean squared error (MSE), root mean squared error (RMSE), cross-entropy and many others. In general MSE, MSE are commonly used for regression problems, with the latter being less popular and cross-entropy also called log loss is used for binary or multi-class classification problems *Janocha* and *Czarnecki*, 2017. Since all problems presented in due course or regression problems, that use MSE as their loss function, this will be the only one emphasized.

$$MSE = \frac{1}{n} \sum_{i=1}^{n} (y_i - \widetilde{y})^2$$
 (4.8)

Equation 4.8 shows the MSE function which is the sum of the squares of the differences of all the predicted and the real values. The same function can be rewritten with the previously used terminology of weights and biases in equation 4.8.

$$L(w,b) = \frac{1}{2n} \sum_{x} ||y(x) - \tilde{y}||^{2}.$$
 (4.9)

With w and b as the collection of all the weights and the biases in the network used to optimize the function y(x). Giving the quadratic nature of the function the L(w,b) will always be positive. And if  $L(w,b) \to 0$  the loss is minimal, meaning that the real and predicted values are close together and the network found weights and biases that explain the output well.

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A widely used function to find the optimum for such a loss function is gradient descent. Its objective is to fine the minimum for the loss function *Bottou*, 1991. The behind gradient descent or other optimizing algorithms is start with randomly initialized weights and biases and repeatedly move them in direction  $\Delta w$  and  $\Delta b$ . This results in a change of the loss function as shown in equation 4.10, making use of partial derivatives.

$$\Delta L = \frac{\partial L}{\partial w} \Delta w + \frac{\partial L}{\partial h} \Delta b \tag{4.10}$$

Ideally  $\Delta L$  is negative and the optimization algorithm found  $\Delta w$  and  $\Delta b$  that lead to a reduction of the loss. To simplify this problem let  $\Delta d$  be the vector of changes:  $\Delta d = (\Delta w, \Delta b)^T$  and  $\nabla L$  the vector of the partial derivatives: equation 4.11

$$\nabla L = \left(\frac{\partial L}{\partial w}, \frac{\partial L}{\partial w}\right)^T \tag{4.11}$$

Having defined  $\nabla L$  and  $\Delta d$  the term 4.10 can be simplified as equation 4.12

$$\Delta C = \nabla L * \Delta d \tag{4.12}$$

Now the task of gradient descent or any other optimizer is to find  $\Delta d$  that results in  $\Delta C$  being negative as shown in equation 4.13

$$\Delta d = -\eta \nabla L \tag{4.13}$$

In this case  $\eta$  is a small positive decimal number, commonly referred to as the learning rate, which usually, but not exclusively ranges from 0.1 to 0.001. However it can be larger or much smaller in some cases. Having found a way to ensure that  $\Delta L$  always decreases according to equation 4.13 it is utilized to repeatedly update the gradient  $\nabla L$ . To make the gradient descent algorithm efficient the learning rate  $\eta$  must be chosen correctly. If  $\eta$  is too large, the gradient  $\Delta L$  might end up being larger than zero, leading to an increase of the loss, and if the step size is too small

convergence will either take too long or not take place at all Bergstra et al., 2011. In practical machine learning approaches different learning rates are tested. There are also algorithmic approaches. While equation 4.10 only accounts for two inputs features, it can be generalized to compute n inputs shown in equation 4.14.

$$\nabla L = \left(\frac{\partial L}{\partial w_1}, \dots, \frac{\partial L}{\partial w_n}\right)^T \tag{4.14}$$

Equation 4.15 shows the gradient descent how it is used to repetitively update the weights and biases to optimize the loss function L(c, w) with w and b as the weight and bias matrices and the learning rate  $\eta$ . In machine learning each iterational update of the network is often called epoch or training epoch.

$$w = w_i - \eta \frac{\partial}{\partial w} L(w) \tag{4.15a}$$

$$b = b_i - \eta \frac{\partial}{\partial h} L(b) \tag{4.15b}$$

(4.15c)

Substituting the partial differentials with  $\nabla L$  equation 4.15 a simplifies to:

$$w = w_i - \eta \nabla L \tag{4.16}$$

#### **Optimizers**

The previous section introduced the concept of gradient descent, an algorithm to minimize the loss function of the weights and biases of a neural network. All other optimizers introduced here, are either variations or extensions of the basic gradient descent algorithm (GD) shown in 4.15. One disadvantage of gradient descent is that

if the data sets grow larger, the demand in memory for computation increases exponentially. Taking into consideration machine learning is a popular method ind big data applications this is a serious drawback. Methods to overcome that are stochastic gradient descent and mini-batch gradient descent. The idea behind the latter is to randomly divide the entity of the training data in sub-samples called mini-batches Bottou and Bousquet, 2008. The network is then trained iteratively over the mini batches. The batch size influences the accuracy and the training speed and is another hyperparameter which has to be tuned. If the batch size is 1 mini batch GD is also referred to as stochastical gradient descent (SGD). During the optimization process optimizers can find local minima in the cost function without being able to overcome them to find the desired global minimum. An algorithm extending GD to accelerate the search of the global minimum is momentum. Which allows the GD to speed up when the loss is decreasing and to slow down when going in the wrong direction - increasing the loss function L(w, b). This is achieved by accounting for the gradient of the previous step in the calculation of the current step. This concept was introduced by *Polyak*, 1964 and re-popularized alongside backpropagation learning by Rumelhart, Hinton, and Williams, 1988.

$$w = w_i - \eta \nabla L + \alpha \Delta w \tag{4.17}$$

Equation 4.17 shows how the momentum is mathematically represented in GD to update the weights w or likewise the biases the delta of the weights multiplied by the coefficient  $\alpha$  - the momentum, which usually ranges from 0.1 to 0.9 and is another parameter to tuned for successful training. If the momentum is two small the GD will not be able to overcome local minima and if  $\alpha$  is two large the loss functions tends to oscillate without finding an optimum LeCun, Bengio, and Hinton, 2015. For both of the momentum and the learning rate it is impractical to remain on the same level during all training epochs. Because after each epoch the loss function is either closer or further away from its global minima and depending on the distance to that

minimum it is desirable to have larger or smaller learning rates and momenta. This can be achieved with naive approaches for example using a step function to gradually decrease those values after each iteration, or to utilize algorithmic approaches *Michie, Spiegelhalter*, and *Taylor*, 1994. There is a large variety of optimizers trying to find optimal values for  $\alpha$  and  $\eta$  and till today this field is under active research *Goodfellow, Bengio*, and *Courville*, 2016. Popular among those are: RMSprop *Hinton*, *Srivastava*, and *Swersky*, 2012; Nesterov momentum *Dozat*, 2016; Adadelta *Zeiler*, 2012; Adagrad *Ruder*, 2016 and Adam *Kingma* and *Ba*, 2014. With Adam being the most widely used optimizer today. Nesterov momentum is slight change to the normal momentum capable of having huge impacts in practical applications, because it helps avoiding oscillations around the minimum by using intermediate information to adapt the momentum.

RMSProp - root mean square propagation - is a method aiming to adapt the learning rate algorithmically, by choosing  $\eta$  for each parameter. And lastly the wide-spread Adam optimizer combines both of the features of momentum and RMSProp and adapts the learning rate as well as the momentum iteratively *Kingma* and *Ba*, 2014.

#### **Backpropagation**

maybe i will leave out backpropagation  $\odot$ 

Backpropagation Rumelhart, Hinton, and Williams, 1988

#### Regularization parameters

When applying the combined aforementioned algorithms and optimizers to find global minima of a loss function of a neural network a problem arises, because optimizers like Adam work "too" well. This issues is due to the fact that neural networks have 100s of thousand of free parameters to be trained, deep neural networks have billions and trillions of parameters. If training of the neural net continues for enough epochs eventually. The loss function will approach a minimum and as  $L(w,b) \to 0$  the initial conclusion could be that the training was quite successful,

but when trying to apply the network trained on the training data set (TRN) to a testing data set (TST). The loss and accuracy of the prediction of TST and TST are very large or accordingly small. This phenomenon is know as overfitting and a lot of fine tuning of hyperparameters is devoted to minimizing this effect *Tetko*, *Living-stone*, and *Luik*, 1995. Figure 4.4 visualizes the effects of overfitting during training *Goodfellow*, *Bengio*, and *Courville*, 2016.

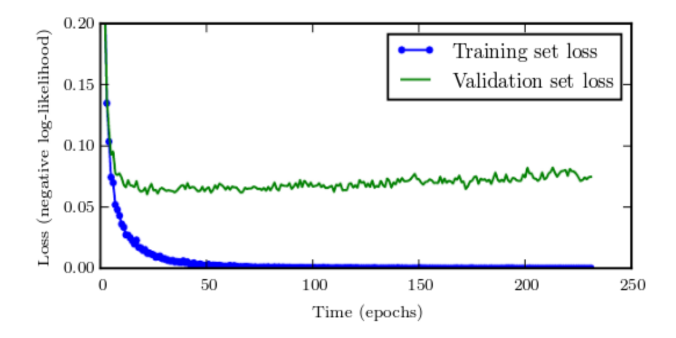


FIGURE 4.4: Learning curves showing how a loss function changes during training in the loss and validation data set. While the training loss approaches 0 the validation loss starts increasing after hitting a minimum. This effect is due to overfitting on the training data set. Figure from *Goodfellow*, *Bengio*, and *Courville*, 2016.

#### **Cross-validation**

A method that is used in basically every training of neural network is splitting up the data sets in multiple sub-sets. More specifically a training set (TRN) and a testing set (TST). The training set is used to minimize the loss functions and its success is evaluated on the TRN set, by comparing the predicted values  $\hat{y}$  with the

real values in TST y. For all neural nets in this study person's correlation coefficient was chosen as performance metric, as in equation 4.18 *Soper* et al., 1917.

$$\rho(y,\hat{y}) = \frac{cov(y,\hat{y})}{\sigma_y \sigma_{\hat{y}}} \tag{4.18}$$

There are other popular performance metrics, especially for classification problems, like AUC (area under the curve) and ROC (receiver operating characteristics), which basically evaluate by weighing sensitivity and specificity. In cross-validation compared to single validation the initial data set is split into TRN and TST multiple times e.g. if the ratio is 80:20 5 times, and each TRN-TST pair is evaluated individually. Sometimes it becomes necessary to use a third subset - the validation data. Because hyperparameter tuning is performed with the TRN and TST sets, a third portion of the data needs to be assessed to check whether the neural network is able to generalize on global data.

L1 and L2 loss

L1 and L2

**Dropout** 

## 4.1.2 On the nature of quantitative traits

According to the omnigenic model which is an extension of the polygenic model proposed by *Boyle*, *Li*, and *Pritchard*, 2017 and thoroughly reviewed in *Timpson* et al., 2018 all traits or phenotypic values are influenced by a great number or all genes in the genome. Therefore resulting in traits following certain gradual statistical distributions instead of being binned in classes or even binary. Intuitively this might be contradicting with the foundation of modern Genetics - Mendel's three laws. That where derived from observations with where mainly influenced by one locus. But staying with one of Mendel's examples the round or wrinkled surfaces of peas

Pisum sativum, an assessment of a couple of thousands peas, would most likely inevitably lead to the conclusion that form the "roundest" to the "wrinkliest" pea any gradual step between those is possible and observable. Mendel's third law of independent segregation also only holds true under certain assumptions. The most simplest one being that the traits under investigation have to be located on different linkage groups. Otherwise for the 7 traits used in Mendel's initial studies would not have segregated independently. The odds of 7 randomly selected traits being on 7 different linkage groups are rather small, especially taking into account, that the genome of the *P. sativum* consists of only 7 chromosomes itself *Kalo* et al., 2004. Mendel probably new about traits not following its own laws, as well as being aware of the quantitative nature of traits such as the constitution of surfaces of peas or the color of petals. But being the pioneer of a then rather unexplored field of science, some of which big questions we fail to satisfactory answer today, he did not have the resources or the knowledge to explain behavior's not "mendeling", that were only able to be deciphered in later decades and centuries based on his ground-breaking work.

Initially thought to be contradicting to Mendel's ideas Darwin proposed the concept's of evolution due to natural selection which also introduce the idea of traits following a gradual distribution *Darwin*, 1859. This contrast led to a long lasting debate in the scientific community in the early 1900s, between the Mendelians and the biometricians who believed in the quantitative nature of continuous traits. This conflict has eventually been solved by Fisher's fundamental work published in 1918 *Fisher*, 1919. His theories combined the then in all fields of science popular research of distributions with genomics. He he mathematically proved that traits influenced by many genes, with randomly-sampled alleles follow a continuous normal distribution in a population. While this combined the ideas of Mendel and the biometricians it opened an other long debated question of effect size and the overall architecture of complex traits. While in the theory of monogenic traits the effect size of the single gene on the trait is 1 or 100 % with an increasing number of genes

influencing a complex traits the *per sè* contribution of single gene has to decrease with an increasing number of loci determining the value a given trait. In the 1990s it has been thought, that complex traits are predominantly controlled from few genes with a large to medium effect size, while others had a minimal influence *Zhang* et al., 2018.

With the upcoming popularity of GWAS as the favored method to decipher genetic architectures of traits, or having pioneered in human genetics in became clear that the majority of the effect sizes are tiny < 1 % while there are very few loci which have a moderate effect on the phenotypic variance of a population with around 10 % or less *Korte* and *Farlow*, 2013, *Stringer* et al., 2011. This nature of quantitative traits present great challenges to animal *Goddard* and *Hayes*, 2009 and plant breeding *Würschum*, 2012, in further improving crop or livestock performances, as well complicating the decomposition of genomic causes for diseases like schizophrenia or autism in human medicine *De Rubeis* et al., 2014, *Purcell* et al., 2014.

While the complex nature of the architecture of quantitative traits provide enough challenges as is, all traits will also be influenced by the environment from which an individual originates. Therefore the distribution of trait values in a given population can be expressed as the addition of the variances of its genetic and the environmental effects 4.19.

$$\sigma_P = \sigma_G + \sigma_E \tag{4.19}$$

The genomic and the environmental effects not only influence the phenotypic variance directly, but the environment also has an influence on gene expression methylation of DNA bases etc. and therefore the equation 4.19 needs to be extend by the variance of the gene-environment interactions  $\sigma_{GxE}$  4.20 , *Lynch* and *Walsh*, 1998, *Walsh* and *Lynch*, 2018b.

$$\sigma_P = \sigma_G + \sigma_E + \sigma_{GxE} \tag{4.20}$$

Equation 4.20 shows the decomposition of the phenotypic variance, to thoroughly understand complex genetic architectures of traits the genetic variance needs to be decomposed further in its additive, dominance and epistatic components 4.21

$$\sigma_G = \sigma_A + \sigma_D + \sigma_I \tag{4.21}$$

The additive effects are caused by single, for this model mostly homozygous, loci while the variance caused by dominance effects, is caused by heterozygous loci and their resulting interactions being full-, over-, co- or underdominant. And lastly the interaction effects that are a result of two or more genes only having an impact if the involved genes co-occur in a certain state. The resulting variance is commonly known as gene-gene interactions and/or epistasis *Falconer* and *Mackay*, 1996. Since possible interactions in a genome can happen between additive or dominant or a combination of those loci. The variance due to interaction effects  $\sigma_I$  can be further dissembled in the variance resulting from additive-additive  $\sigma_{AA}$  dominant-dominant  $\sigma_{DD}$  and additive-dominant  $\sigma_{AD}$  terms as represented in equation 4.22.

$$\sigma_I = \sigma_{AxA} + \sigma_{DxD} + \sigma_{AxD} \tag{4.22}$$

Knowledge of the variance components involved in the expression of a trait in population, lead up to the estimation of the total influence of all genetic variances and the environmental variance one the phenotypic distribution. This concept is called heritability. The heritability of a trait  $H^2$  accounts for the proportion of the phenotypic variance controlled by the total genetic variance as shown in equation 4.23. This is also referred to as broad sense heritability, because all genetic effects including additive, dominance and epistatic effects are included *Brooker*, 1999.

$$H^2 = \frac{\sigma_A + \sigma_D + \sigma_I}{\sigma_P} \tag{4.23}$$

The concept of narrow-sense heritability 4.24 is similar to the broad-sense heritability, but only the additive genetic effects are included in the genetic part of the equation. This differentiation is import for natural and artificial selection and thus is commonly used in evolutionary genomics and breeding. Because in diploid species each parent only passes down on a single a allele of a given locus, dominance effects or interaction effects are not commonly inherited from one parent. Therefore it is mainly the additive genetic effects of a parent that influences its offspring. While the dominance and epistatic variances are controlled by the combination of the parents *Falconer* and *Mackay*, 1996, *Walsh* and *Lynch*, 2018b.

$$h^2 = \frac{\sigma_A}{\sigma_P} \tag{4.24}$$

# 4.1.3 Artificial selection in plant and animal breeding in the genomics era

#### **Introduction to genomic selection**

Genomic prediction has been applied to almost all relevant crop and model species. Including: A.thaliana Hu et al., 2015; Shen et al., 2013. Alfalfa (Medicago sativa) Li and Brummer, 2012; Annicchiarico et al., 2015; Li et al., 2015; Biazzi et al., 2017; Hawkins and Yu, 2018. Barley Neyhart, Lorenz, and Smith, 2019; Oakey et al., 2016; Zhong et al., 2009. Cassava (Manihot esculenta) Elias et al., 2018a; Elias et al., 2018b. Cauliflower (Brassica olearacea spp) Thorwarth, Yousef, and Schmid, 2018. Cotton (Gossiypium spp.) Gapare et al., 2018. Maze (Zea mays) Moeinizade et al., 2019; Allier et al., 2019; Brauner et al., 2018; Schrag et al., 2018; Schopp et al., 2017b; Sousa et al., 2017; Schopp et al., 2017a; Kadam et al., 2016; Bustos-Korts et al., 2016a; Montesinos-López et al., 2015; Owens et al., 2014; Lehermeier et al., 2014; Technow et al., 2014; Peiffer et al., 2014; Riedelsheimer et al., 2013; Guo et al., 2013; Technow, Bürger, and Melchinger, 2013; Windhausen et al., 2012; Rincent et al., 2012. Potato (Solanum tuberosum); Enciso-Rodriguez et al., 2018; Endelman et al., 2018. Rape seed (Brassica naps) Würschum, Abel, and Zhao, 2014; Jan

et al., 2016; Luo et al., 2017; Werner et al., 2018; Snowdon and Iniguez Luy, 2012; Qian, Qian, and Snowdon, 2014. Rice (Oryza sativa) Momen et al., 2019; Hassen et al., 2018; Xu, 2013; Grenier et al., 2015. Rye (Secale cerale) Auinger et al., 2016; Bernal-Vasquez et al., 2014; Wang et al., 2014; Bernal-Vasquez et al., 2017; Marulanda et al., 2016. Sugar beet (Beta vulgaris), Würschum et al., 2013; Biscarini et al., 2014. Sugar cane (Saccharum officinarum) Gouy et al., 2013 Soybean (Glycine max) Stewart-Brown et al., 2019; Jarquin, Specht, and Lorenz, 2016; Xavier, Muir, and Rainey, 2016. Switchgrass (Panicum virgatum) Poudel et al., 2019; Ramstein and Casler, 2019; Ramstein et al., 2016. Wheat (*Triticum aestivum*) Cuevas et al., 2019a; Howard et al., 2019; Krause et al., 2019; Rincent et al., 2018; Norman et al., 2018; Belamkar et al., 2018; Ovenden et al., 2018; Sukumaran et al., 2016; Bustos-Korts et al., 2016b; Gianola et al., 2016; Crossa et al., 2016; Thavamanikumar, Dolferus, and Thumma, 2015; Lopez-Cruz et al., 2015. As well as various tree species *Almeida Filho* et al., 2019; *Rincent* et al., 2018; *Kainer* et al., 2018; Ratcliffe et al., 2017; El-Dien et al., 2016; Kumar et al., 2015; Jaramillo-Correa et al., 2014; Zapata-Valenzuela et al., 2013; Holliday, Wang, and Aitken, 2012; Resende et al., 2012. Even though GS finds broad application in plant breeding it has been originally developed for the use in animal breeding Hayes and Goddard, 2010; Goddard, Hayes, and Meuwissen, 2011. The gold standard is a method known as genomic BLUP VanRaden, 2008 which utilizes a relationship matrix based on the co-occurrence of genetic markers. This method is derived from the pre-genomic era in animal breeding where the relationship matrix was constructed after pedigrees according to the best linear unbiased predictors based linear mixed models developed by *Henderson*, 1975. GBLUP accounts only for additive-genetic effects VanRaden, 2008. There are other methods that are able to account for more complex genomic effects that are non-additive. Popular among those are reproducing Kernel Hilbert Spaces (RKHS) Gianola and Kaam, 2008. Alternatively to linear mixed models a variety of different Bayesian methods became popular, basically differing in the degree of shrinkage of the assumed marker distribution *Hayes* and *Goddard*, 2001; *Gianola* et al., 2009; *Habier* et al., 2011; Gianola, 2013; Crossa et al., 2017; Azodi et al., 2019.

#### Genomic selection in recurrent selection and the breeders equation

While the quantitative genetic methods breeders utilize are complex their goals can be defined in one sentence: To genetically improve plant germplasms for agriculture. The breeding process started at the same time as farming was first practiced 10.000 BC in the region between the Euphrat and Tigris rivers known as the fertile crescent. This changed the phenotypic appearance of the early crops dramatically to the point that they share little external traits with their wild ancestors. Those changes have also been deeply carved into the genomes, that underwent serious alterations, including hybridization, duplications etc. Leading to most crop plants not having any wild ancestors with whom they could naturally mate. For example wheat (Triticum aestivum), one of the three most important sources of food on a global scale underwent multiple hybridization steps *Ozkan*, *Levy*, and *Feldman*, 2001. Wheat is a hybrid from either the diplod emmer (*T. diccoides*) or durum wheat (*T. du*rum) and Aegilops tauschii, while emmer and durum are hybrids derived from wild emmer which is a hybrid of wild grass of the genus of Aegilops and T. urata Friebe et al., 2000; Feldman and Levy, 2012. While being ignorant of modern genetics early "plant breeders" must have had an intuitive, yet naive, understanding of the general concept of heritability in a way that they must have established that offsprings share properties of their parent generation, which lead to regrowing individuals with desired traits generations after generation. This lead to many changes including that artificial selected plants are commonly largely inbred. This process could be considered an early form of recurrent truncation selection. Truncation selection on a normal distributed phenotype is shown in figure 4.5.

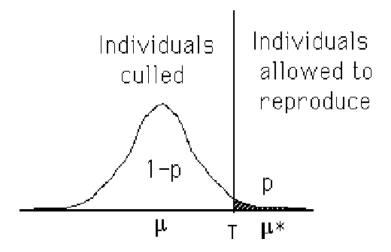


FIGURE 4.5: Truncation selection from a normal distributed phenotype with a threshold value of T,  $\mu$  as the mean of the total population and  $\mu^*$  as the mean of the selected phenotypes. Graphic from *Walsh* and *Lynch*, 2018a

Like the early breeders modern breeders of to determine the selection threshold T to divide the total population with the mean  $\mu$  into two groups: the individuals culled and the ones allowed to reproduce with the mean  $\mu^*$ . The difference between those two is the selection differential S:

$$S = \mu^* - \mu \tag{4.25}$$

which in the case of normal distributed data as depicted in figure ?? can be expressed as:

$$S = \varphi(\frac{T - \mu}{\sigma})\frac{\sigma}{p} \tag{4.26}$$

From which we can obtain the selection intensity i, which makes i solely a function of p

$$i = \frac{S}{\sigma} = \frac{\varphi(z_{|1-p|})}{p} \tag{4.27}$$

With recurrent truncation selection over many generations the population mean of the trait  $\mu$  will change (hopefully in the desired direction), if the heritability (in

this case the narrow sense heritability)  $h^2 > 0$ . It is impossible to breed for traits that do not contain any genetic components in its architecture *Walsh* and *Lynch*, 2018b. Next to *i* the intensity and  $h^2$  the additive portion of the heritability the accuracy of the selection  $r_{uA}$  is important for the success of a breeding program. Those three terms can be applied to compute the gain of selection *R* over one generation (equation 4.28). Due to its importance in the evaluation of breeding schemes it is known as the breeder's equation *Mousseau* and *Roff*, 1987; *Falconer* and *Mackay*, 1996; *Kingsolver* et al., 2001.

$$R = i r_{uA} \sigma_A \tag{4.28}$$

The accuracy  $r_{uA}$  of equation 4.28 in cases only phenotypic selection is conducted the heritability and in cases where the selection process is aided by genomic prediction it is the prediction accuracy. According to breeder's equation there are three parameters which can be influenced through genomic prediction.

(i) The prediction accuracy, which is usually smaller than the heritability, varies for different prediction equations and an increase in the accuracy will lead to an proportional increase in R the gain per generation cycle. For this reasons since 2001 in quantitative genetics one very active field of research was and still is to find new algorithms that are superior to others as presented in the next chapter (4.1.3. As later evaluated on more than 150 phenotypes in chapter ??  $h^2$  is always larger than  $r_{uA}$ . Which if it was the only variable factor in equation 4.28 make genomic selection inferior to phenotypic selection. Which from a certain point of view it is. Phenotypic trials are better approximation for phenotypic appearance as GEBVs. However as the cost of genotyping have decreased dramatically in the last 20 years. Phenotyping in the field remains tedious, laborious and mostly vastly expensive. Taking into account that field trials have to be repeated in several years and locations to produce reliable accounts it becomes clear that genotyping 10th of thousands of accessions is much cheaper than conducting field trials with 1000 of them.

(ii) The selection intensity can much stricter if the total population that is selected from is larger and in genomic prediction settings they are, because it allows selection from two pools. The first the pool of plants with known phenotyped and known genotype and from those were just genomic data is available. When selecting from a pool of 1000 with p=0.05 with the goal to keep 50 plants in the next breeding cycle, the same goal can be reached when genomically selecting from a pool of 10000 with and intensity of p=0.05.

(iii) The decreased time per generation is probably the largest advantage of genomic selection when applied to breeding. While in field trials it is only possible to have one generation per year. Genomic selection does not require the plants to be grown in the field. For selection it is only necessary to grow enough so that DNA can be extracted from the tissue and evaluated. After selection only the ones above the threshold are grown until they bear seats (or any other reproductive organ) and be used for the next selection cycle, allowing up to ten generations per year. This development hast lead to the rise to a new branch of breeding: speed breeding *Ghosh* et al., 2018; *Watson* et al., 2018. In practical, company-level genomic prediction as largely contributed to an increase by a factor of 2 of the gain in selection in recent years (Viktor personal communication).

The last term in equation 4.28 the additive genetic variance  $\sigma_A$  is not directly yet heavily influenced by the described breeding scheme. Artificial selection has similar effects on the genetic variance as bottlenecks in natural selection have: it decreases, thus making it harder to increase R in later selection cycles Walsh and Lynch, 2018b.

#### Genomic BLUP and Bayesian methods

All methods share a common statistical obstacle which is commonly referred to as the n >> p problematic, which arises because the number n marker is usually significantly larger than the number of observations p. In practical applications it is not uncommon the have more than 100k markers while the number of phenotypes is no larger than 100. This does not allow to obtain genomic estimated breeding values

(GEBV) by single marker regression as done by GWAS, which have highly inflated over all SNP-effects *Korte* and *Farlow*, 2013. One possible is to include effect sizes as random effects and make prior assumptions about their distribution. The difference in prior distribution is the main distinction between the methods of the Bayesian alphabet *Gianola*, 2013.

#### **Genomic BLUP**

In the early years of research on genomic prediction algorithms they were not solely benchmarked against each other, but had to compete with the previously popular pedigree methods. Quickly in the course of the first decade of this millennium the superiority of the genomic method became elucidated in livestock and plant breeding Habier, Fernando, and Dekkers, 2007; VanRaden, 2008; VanRaden et al., 2008; Harris, Johnson, and Spelman, 2009. While the genomic methods are superior to nongenomic methods, there is no clear evidence that either of the genomic methods are suerior to each other and there is lack of empirical evidence that the Bayesian methods generally outperform GBLUP Moser et al., 2009; Bernardo, 2010; Azodi et al., 2019. Alike in the pedigree BLUP in genomic BLUP the covariance between related individuals is used for the prediction. In the later case it is constructed from marker information. In the GWAS terminology the relationship matrix is referred to as *K* for kinship in GWAS literature, while in GS circumstances it is also called GRM (genomic relationship matrix) or simply abbreviated as G. This study will remain consistent with the circumstantial literature and therefore purposely inconsistent within itself. In the chapter addressing GWAS the it will be called *K* for kinship matrix and it the following chapter elucidating GBLUP it will be referred to as G. The general genomic prediction model 4.29 is derived from the mixed model *Henderson*, 1975; VanRaden, 2008 and implemented as:

$$Y = X\beta + Zu + \varepsilon \tag{4.29}$$

where Y is and nx1 vector of phenotypic observations, X the matrix of the fixed effects and  $\beta$  the vector of the fixed effects. Z the incidence matrix for the combined marker effects and u a (nx1) vector of the additive genetic effect the vector of the residuals  $\varepsilon$ . To construct the model lets assume a matrix of size (nxm) with n individuals and m loci M containing marker information for 3 individuals on 4 loci, thus being of size 3x4. The 4 markers of matrix 4.30 can take values of -1, 0 and 1 translating into minor allele, heterozygous locus and major allele. The following example calculation has been adapted from Isik, 2013.

$$M = \begin{pmatrix} -1 & 0 & 1 & -1 \\ -1 & 0 & 0 & 0 \\ 0 & 1 & 1 & -1 \end{pmatrix} \tag{4.30}$$

The M matrix contains all the information that is necessary for the computation of the K matrix and other viable genetic parameters. The MM' matrix of size nxn (??) bears additional parameters.

$$MM' = \begin{pmatrix} 3 & 1 & 2 \\ -1 & 1 & 0 \\ 2 & 0 & 3 \end{pmatrix} \tag{4.31}$$

The diagonal show the number of homozygous loci per individual, while the other elements of the matrix indicate the number of markers shared by related individuals and is an indicator for the distance of the relationship as defined by identity-by-descent VanRaden, 2008; Misztal et al., 2013. While matrix 4.31 calculates the metrics per individual the M'M matrix (4.32) counts those metrics per marker. Likewise the diagonal contains the number of homozygous individuals per marker.

$$M'M = \begin{pmatrix} 3 & -1 & 0 & 0 \\ -1 & 1 & 1 & 1 \\ 0 & 1 & 2 & 1 \\ 0 & 1 & 1 & 2 \end{pmatrix}$$
(4.32)

The next step is to obtain a matrix of the allele frequencies at each locus of the size (nxm) as matrix M. For the design of matrix P (4.33) let the minor allele frequencies  $p_1 \dots p_4$  be {0.3, 0.2, 0.1, 0.15}. The allele frequency of the  $i^{th}$  column of P is expressed, according the  $i^{th}$  marker of matrix M as  $P_i = 2(p_i - 0.5)$  resulting in:

$$P = \begin{pmatrix} -0.4 & -0.6 & -0.8 & -0.7 \\ -0.4 & -0.6 & -0.8 & -0.7 \\ -0.4 & -0.6 & -0.8 & -0.7 \end{pmatrix}$$
(4.33)

The allele frequencies as in this simulated example should be taken from the entire population and not the subsample used in this calculation VanRaden, 2008. The final step to obtain the Z matrix of equation 4.29 is to subtract the P matrix from the M matrix Z = M - P resulting in:

$$Z = \begin{pmatrix} 1.4 & 0.6 & 1.8 & -0.3 \\ -0.6 & 0.6 & 0.8 & 0.7 \\ 0.4 & 1.6 & 1.8 & -0.3 \end{pmatrix}$$
(4.34)

In Z the mean values of the allele effects are set to 0 and the subtraction of P emphasizes the effect of rare variants VanRaden, 2008. There is a large variety of methods to generate the genomic relationship matrices and here lies the major difference between different genomic BLUP methods. But they all have in common that K is always of size nxn.

(i) The naive is approach to iterate over each individual and count the common markers with every other individual. This approach is not uncommon and suited for inbred or doubled-haploid populations, less so for outcrossed populations with

high degrees of heterozygosity because in the sample implementation it does only account for homozygous loci. This method becomes computationally intense when the data sets grow larger as common today (personal observation).

(ii) Probably the most popular method in GS is to obtain *K* as proposed by *Van-Raden*, 2008 designed after Wright's *Wright*, 1922 equations for the covariance in structured populations, as described by equation 4.35 with *Z* as in 4.34.

$$G = \frac{ZZ'}{2\Sigma p_i(1-p_i)} \tag{4.35}$$

(iii) The unified additive relationship  $G_{UAR}$  according to Yang et al., 2010 and equation 4.36

$$G_{UAR} = A_{jk} = \frac{1}{N} \Sigma_i A_{ijk} = \begin{cases} \frac{1}{N} \Sigma_i \frac{(x_{ij} - 2p_i)(x_{ik} - 2p_i)}{2p_i(1 - p_i)}, j \neq k \\ 1 + \frac{1}{N} \Sigma_i \frac{x_{ij}^2(1 + 2p_i)x_{ij} + 2p_i^2}{2p_i(1 - p_i)}, j = k \end{cases}$$
(4.36)

where  $p_i$  is the allele frequency at locus i and  $x_{ij}$  the genotype for the  $j^{th}$  individual at the  $i^th$  locus. Another method also proposed by Yang et al., 2010 is to adjust  $G_{UAR}$  with  $\beta$  as in equation ??

$$G_{UARadj} = \begin{cases} \beta A_{jk}, & j \neq k \\ 1 + \beta (A_{jk} - 1), & j = k \end{cases}$$
 (4.37)

(iv) Another approach is to weigh marker by the reciprocals of their expected variance according to the model 4.38 originally designed to investigate population structures in human data *Leutenegger* et al., 2003; *Amin, Van Duijn*, and *Aulchenko*, 2007.

$$G = ZDZ', with$$

$$D_{ii} = \frac{1}{m|2p_i(1-p_i)|}$$
(4.38)

(v) Other methods like the gaussian kernel compute kinship between individuals by the euclidean distance between the respective genotypes *Morota* and *Gianola*, 2014.

$$K(x_{i}, x_{j}) = exp(-\theta d_{ij}^{2})$$

$$= \prod_{k=1}^{m} exp(-\theta (x_{ik} - x_{jk})^{2})$$
(4.39)

with  $d_{ij} = \sqrt{(x_{i1} - x_{j1})^2 + \dots + (x_{ik} - x_{jk})^2 + \dots + (x_{im} - x_{j,a})^2}$  and  $x_{ik}(i, j = 1, \dots, n, k = 1, \dots, m)$  and  $x_{ik}$  as the  $i^{th}$  individual at SNP k.

The linear model in equation 4.29  $Y = X\beta + Zu + \varepsilon$  with  $\beta$  as the vector fixed effects and u as the vector of additive genetic effects. The mixed model can be solved to obtain genomic estimated breeding values as:

$$\begin{pmatrix} X'X & X'Z & 0 \\ Z'X & Z'Z + G^{11} & G^{12} \\ 0 & G^{21} & G^{22} \end{pmatrix} \begin{pmatrix} \hat{b} \\ \hat{y}_1 \\ \hat{y}_2 \end{pmatrix} = \begin{pmatrix} X'y \\ Z'y \\ 0 \end{pmatrix}$$
(4.40)

with  $G^{12}$  as the part of  $G^{-1}$  containing individuals <u>with</u> phenotypic data. with  $G^{22}$  as the part of  $G^{-1}$  containing individuals <u>without</u> phenotypic data and just SNP information.

The GEBV of the unknown phenotypes  $\hat{y}_2$  can thus be predicted as:

$$\hat{y}_2 = -\left(G^{22}\right)^{-1} G^{21} \hat{y}_1 \tag{4.41}$$

GBLUP is fairly easy compared to more complex Bayesian methods and be quickly implemented in any programming language capable of solving liner equations. Computational as the number of phenotypes in the study increases in numbers the timed demand grows exponentially, because the kinship matrix quadruples in size and it becomes more complicated to compute the inverse of *G* (personal observations).

#### **Bayesian methods**

Next to the universal GBLUP a set of related algorithms became popular for solving the mixed models involved in genomic selection, known as the Bayesian alphabet *Gianola* et al., 2009; *Gianola*, 2013. They are all based on Bayes' fundamental theorem in equation 4.42

$$P(\theta|y) = \frac{P(\theta)P(y|\theta)}{P(y)}$$
(4.42)

with  $P(\theta)$  as the prior distribution,  $P(y|\theta)$  as the likelihood and P(y) as the marginal density of y. The prior distribution in GS assume that y was drawn from a certain distribution. Infinitesimal models assume that the genetic effects follow a normal distribution Legarra, Lourenco, and Vitezica, 2018. The Bayesian frameworks however will assume non-normal distributed marker effects. This can be explained by a two-step hierarchical distribution. Stage one assumes that every marker has a priori a different variance Legarra, Lourenco, and Vitezica, 2018.

$$p(a_i|\sigma_{ai}^2) = N(0, \sigma_{ai}^1) \tag{4.43}$$

The second stage assumes prior distributions for the variances.

$$p(a_i|variable) = P(\dots) \tag{4.44}$$

with *variable* standing for the large variety of prior distribution. In total there are more than >20 different Bayesian models known to the authors with unknown number of different methods proposed. Their main difference "simply" lies in the a priori assumptions. This change can make some methods mathematically much more advanced then others. And as shown in later chapters none of the methods is completely superior over others in terms of prediction accuracy. Approximation to

the solution of the linear equations is usually performed by Gipp's sampling using Markov Chain Monte Carlo (MCMC) simulations *De Los Campos* et al., 2009; *Campos* and *Rodriguez*, 2016. Table 4.1 summarizes commonly applied Bayesian methods for genomic prediction indicating the key differences between them.

TABLE 4.1: Overview of properties of a variety of commonly applied Bayesian methods for genomic prediction. Table altered after *Kärkkäinen* and *Sillanpää*, 2012

Name	Reference	Prior	Indicator	Hierarchy	Hyperprior	Estimation
BayesA	Hayes and Goddard, 2001	Student	No	Yes	No	MCMC
BayesB	Hayes and Goddard, 2001	Student	Yes	Yes	No	MCMC
BayesC	<i>Verbyla</i> et al., 2009	Student	Yes	Yes	No	MCMC
BL	Xu, <mark>2010</mark>	Laplace	No	Yes	No	EM
BayesD $\pi$	Habier et al., 2011	Student	Yes	Yes	Yes	MCMC

The name is given by the author. The prior column tells which shrinkage prior is used.

#### cross validation

## 4.1.4 Genomic selection using artificial neural networks

Genomic selection (GS) has been successfully applied in animal *Gianola* and *Rosa*, 2015; *Hayes* and *Goddard*, 2010 and plant breeding *Crossa* et al., 2010; *Desta* and *Ortiz*, 2014; *Heffner* et al., 2010; *Crossa* et al., 2017 as well as in medical applications, since it was first reported *Hayes* and *Goddard*, 2001. Since then the repertoire of methods for predicting phenotypic values has increased rapidly e.g. *De Los Campos* et al., 2009; *Habier* et al., 2011; *Gianola*, 2013; *Crossa* et al., 2017. The most commonly applied methods include GULP and a set of related algorithms known as the bayesian alphabet *Gianola* et al., 2009. Genomic prediction in general has repeatedly been shown to outperform pedigree-based methods *Crossa* et al., 2010; *Albrecht* et al., 2011 and is nowadays used in many plant and animal breeding schemes. It has also

been shown that using whole-genome information is superior to using only featureselected markers with known QTLs for a given trait Bernardo and Yu, 2007; Heffner, Jannink, and Sorrells, 2011 in some cases. A more recent study Azodi et al., 2019 compared 11 different genomic prediction algorithms with a variety of data sets and found contradicting results, indicating that feature selection can be usefull in some cases the when the whole genome regression is performed by neural nets 1 While every new method is a valuable addition to the tool-kits for genomic selection, some fundamental problems remain unsolved, of which the n»p problematic stands out. Usually in genomic selection settings the size of the training population (TRN) with n phenotypes is substantially smaller than the number of markers (p) Fan, Han, and Liu, 2014. Making the number of features immensely large, even when SNP-SNP interactions are not considered. Furthermore each marker is treated as an independent observation neglecting collinearity and linkage disequilibrium (LD). Further difficulties arise through non-additive, epistatic and dominance marker effects. The main problem with epistasis issue quantitative genetics is the almost infinite amount of different marker combinations, that cannot be represented within the size of TRN in the thousands, the same problems arises for example in GWA studies *Korte* and Farlow, 2013. With already large p the number of possible additive SNP-SNP interactions potentiates to  $p^{(p-1)}$ . Methods that attempt to overcome those issues are EG-BLUP, using an enhanced epistatic kinship matrix and reproducing kernel Hilbert space regression (RKHS) Jiang and Reif, 2015; Martini et al., 2017.

In the past 10 years, due to increasing availability of high performance computational hardware with decreasing costs and parallel development of free easy-to-use software, most prominent being googles library TensorFlow *Abadi* et al., 2016 and Keras *Chollet*, 2015, machine learning (ML) has experienced a renaissance. ML is a set of methods and algorithms used widely for regression and classification problems. popular among those are e.g. support vector machines, multi-layer perceptrons (MLP) and convolutional neural networks. The machine learning mimics the

architecture of neural networks and are therefore commonly referred to as artificial neural networks (ANN). Those algorithms have widely been applied in many biological fields *Min*, *Lee*, and *Yoon*, 2017; *Lan* et al., 2018; *Mamoshina* et al., 2016; *Angermueller* et al., 2016; *Webb*, 2018; *Rampasek* and *Goldenberg*, 2016.

A variety of studies assessed the usability of ML in genomic prediction González-Camacho et al., 2018; González-Camacho et al., 2016; Ogutu, Piepho, and Schulz-Streeck, 2011; Montesinos-López et al., 2019a; Grinberg, Orhobor, and King, 2018; Cuevas et al., 2019b; Montesinos-López et al., 2019b; Ma et al., 2017; Qiu et al., 2016; González-Camacho et al., 2012 Li et al., 2018. Through all those studies the common denominator is that there is no such thing as a gold standard for genomic prediction. No single algorithm was able to outperform all the others tested in a single of those studies, let alone in all. While the generally aptitude of ML for genomic selection has been repeatedly shown, how no evidence exists that neural networks can outperform or in many cases perform on that same level as mixed-model approaches as GBLUP Hayes and Goddard, 2001. While in other fields like image classification neural networks have up to 100s of hidden layers *He* et al., 2016 the commonly used fully-connected networks in genomic prediction of 1 - 3 hidden layers. With 1 layer networks often being the most successful among those. Contradicting to the idea behind machine learning in genomic selection 1 hidden layer networks will be inapt to capture interactions between loci and thus only account for additive effects. As shown in Azodi et al., 2019 convolutional networks perform worse than fully-connected networks in genomic selection, which again is contradicting to other fields where convolutional layers are applied successfully, e.g natural language processing Dos Santos and Gatti, 2014 or medical image analysis Litjens et al., 2017. Instead of using convolutional layers and fully-connected layers only, as show in Pook et al 2019, we also propose to use locally-connected layer in combination with fully-connected layers. While CL and LCL are closely related they have a significant difference. While in CL weights are shared between neurons in LCLs each neuron as its own weight. This leads to a reduced number of parameters to be trained in the following FCLs, and

should therefore theoretically lead to a decrease in overfitting a common problem in machine learning. To evaluate the results of Pook et al. 2019 accomplished with simulated data we used the data sets generated in the scope of the 1001 genome project of *Arabidopsis thaliana Alonso-Blanco* et al., 2016

## 4.2 Proof of concept for ANN-based genomic selection

Having established the quantitative architecture of traits in section 4.1.2 and the basics of machine learning and neural nets in section 4.1.1, that knowledge can be used to provide a proof of concept that neural networks are a candidate for GP. Table 4.2 provides also the possible genotypes that can be derived by two bi-allelic markers  $G_1 \dots G_4$  on a fictional haploid organism. In this simulation the effect sizes for each marker  $\beta_1$  and  $\beta_2$  are constant with a value of 1.

TABLE 4.2: Simple simulated phenotypes and genotypes for genomic prediction with genotypes  $G_1 \dots G_4$ ,  $M_1$  and  $M_2$  and phenotypes based on additive effects or *and*, *or*, *xor* logic gates.

	$M_1$	$M_2$	$Y_{ADD}$	$Y_{AND}$	$Y_{OR}$	$Y_{XOR}$
$G_1$	0	0	0	0	0	0
$G_2$	0	1	1	0	1	1
$G_3$	1	0	1		1	1
$G_4$	1	1	2	1	1	0

The four phenotypes  $Y_{ADD}$ ,  $Y_{AND}$ ,  $Y_{OR}$  and  $Y_{XOR}$ , which were derived from their respective marker effects.  $Y_{ADD}$  is a phenotype with purely additive effects. So in the nomenclature introduced in chapter 4.1.2  $\sigma_A = \sigma_G$  and  $\sigma_I = 0$ . Since the hypothetical organism is haploid there are dominance effects to be accounted for  $\sigma_D = 0$ . Since all the genetic effects are caused by additive effects and there are now environmental effects  $\sigma_E$ , the narrow sense heritability  $h^2$  - equation 4.24 - and the broad sense heritability  $H^2$  - equation 4.23 - are equally 1. The other three

phenotypes are base on epistatic effects  $\sigma_I$  generated by passing the markers  $M_1$  and  $M_2$  through their respective logic gates. This theoretically in results in  $h^2=0$  and  $H^2=1$ , because there are no additive effects. For  $y_{AND}$  however  $h\approx 0.5$ , because there is a correlation between  $Y_{ADD}$  and  $Y_{AND}$ . In practical applications this allows methods like GBLUP, designed to account for additive genetic effects to capture some of the epistatic effects of  $\sigma_I$  *Vieira* et al., 2017.

According to chapter 4.1.1 a single perceptron would fail to solve *xor* gates. While a network with multiple nodes and layers should be able to overcome that deficit. A relatively simple neural network with two fully-connected hidden layers with 10 and 5 nodes, was trained for the prediction of each phenotypes. To keep the simulation as possible, no regularization parameters, dropout etc. was included. The activation function was ReLU (4.6) with an Adam optimizer. The results of the prediction are shown in table 4.3.

TABLE 4.3: Results of genomic prediction from phenotypes and genotypes in table 4.2

	$M_1$	$M_2$	$\hat{Y}_{ADD}$	$\hat{Y}_{AND}$	$\hat{Y}_{OR}$	$\hat{Y}_{XOR}$
$G_1$	0	0	0.01	0.00	0.00	0.01
$G_2$	0	1	0.99	0.00 0.01 0.00	0.99	0.98
$G_3$	1	0	0.99	0.00	0.99	1.01
$G_4$	1	1	1.99	0.98	1.01	0.02

Not surprisingly, the simple network is able to solve all four phenotypes and predicting the phenotypes accurately. The task was rather easy because the training data set and the testing data set were the same, but it served the purpose of showing that neural networks a generally apt to solve different marker interactions. *In natura* those interactions and the overall genetic architecture is much more complex. Effect sizes are not constant and epistasis may be caused be interactions my more than just two markers, and with an increasing number of markers n the number of possible two-way interactions increases even more so to  $2^{n-1}$ . Smaller interaction effects

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could be obscured under larger additive effects, gene-environment might have a significant influence leading to a model that does not converge.

### 4.3 Material

- 4.3.1 DH populations derived from MAZE landraces
- 4.3.2 A. thaliana
- 4.4 Methods
- 4.4.1 ANN
- **4.4.2 GBLUP**
- 4.5 Results
- 4.6 Discussion

# 5 GWAS

5.1	Reevalulation of 463 phenotypes from the AraPheno
	database

- 5.1.1 Introduction
- 5.1.2 Material and Methods
- 5.1.3 Results
- 5.1.4 Results
- 5.1.5 Disucssion
- 5.2 GWAS in DH landrace populatios of maze across and within environments
- 5.2.1 Introduction
- 5.2.2 Material and Methods
- 5.2.3 Results
- 5.2.4 Results
- 5.2.5 Disucssion

# A Source code

# A.1 GBLUP example

```
1 M <- matrix(c(1,-1,0,0,0,1,1,0,1,-1,0,-1),nrow=3)</pre>
2 MtM <- M %*% t(M)
3 MtM
       [,1] [,2] [,3]
5 [1,]
        3 -1
                  2
      -1 1 0
6 [2,]
7 [3,]
        2
8 tMM <- t(M) %*% t(M)</pre>
9 tMM
[,1] [,2] [,3] [,4]
      2 0 1 -1
11 [1,]
12 [2,]
                      - 1
13 [3,] 1 1 2 -2
14 [4,]
      -1 -1 -2 2
16 P = matrix(rep(2*(mafs - 0.5),3),nrow=3,byrow=T)
17 P
     [,1] [,2] [,3] [,4]
19 [1,] -0.4 -0.6 -0.8 -0.7
20 [2,] -0.4 -0.6 -0.8 -0.7
21 [3,] -0.4 -0.6 -0.8 -0.7
```

## A.2 gwas.py

```
import os
import sys
import time
import numpy as np
```

```
5 import pandas as pd
6 import main
7 import h5py
9 # set defaults
10 \text{ mac_min} = 1
11 batch_size = 500000
out_file = "results.csv"
13 m = 'phenotype_value'
14 perm = 1
15 mac_min = 6
17 X_file = 'gwas_sample_data/AT_geno.hdf5'
18 Y_file = 'gwas_sample_data/phenotype.csv'
19 K_file = 'gwas_sample_data/kinship_ibs_binary_mac5.h5py'
for i in range (1,len(sys.argv),2):
      if sys.argv[i] == "-x" or sys.argv[i] == "--genotype":
          X_file = sys.argv[i+1]
      elif sys.argv[i] == "-y" or sys.argv[i] == "--phenotype":
          Y_file = sys.argv[i+1]
      elif sys.argv[i] == "-k" or sys.argv[i] == "--kinship":
28
          K_file = sys.argv[i+1]
29
      elif sys.argv[i] == "-m":
          m = sys.argv[i+1]
      elif sys.argv[i] == "-a" or sys.argv[i] == "--mac_min":
          mac_min = int(sys.argv[i+1])
      elif sys.argv[i] == "-bs" or sys.argv[i] == "--batch-size":
34
          batch_size = int(sys.argv[i+1])
      elif sys.argv[i] == "-p" or sys.argv[i] == "--perm":
36
          perm = int(sys.argv[i+1])
      elif sys.argv[i] == "-o" or sys.argv[i] == "--out":
          out_file = sys.argv[i+1]
39
      elif sys.argv[i] == "-h" or sys.argv[i] == "--help":
          print("-x , --genotype :file containing marker information in
41
     csv or hdf5 format of size")
```

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```
print("-y , --phenotype: file container phenotype information
     in csv format" )
          print("-k , --kinship : file containing kinship matrix of size
      k X k in csv or hdf5 format")
          print("-m : name of columnn containing the phenotype : default
44
      m = phenotype_value")
          print("-a , --mac_min : integer specifying the minimum minor
     allele count necessary for a marker to be included. Default a = 1"
          print("-bs, --batch-size : integer specifying the number of
46
     markers processed at once. Default -bs 500000" )
          print("-p , --perm : single integer specifying the number of
     permutations. Default 1 == no perm ")
          print("-o , --out : name of output file. Default -o results.
48
     csv ")
          print("-h , --help : prints help and command line options")
          quit()
      else:
          print('unknown option ' + str(sys.argv[i]))
          quit()
53
57 print("parsed commandline args")
59 start = time.time()
61 X, K, Y_, markers = main.load_and_prepare_data(X_file, Y_file, K_file, m)
64 ## MAF filterin
65 markers_used , X , macs = main.mac_filter(mac_min,X,markers)
67 ## prepare
68 print("Begin performing GWAS on ", Y_file)
70 if perm == 1:
output = main.gwas(X,K,Y_,batch_size)
```

```
if( X_file.split(".")[-1] == 'csv'):
          chr_pos = np.array(list(map(lambda x : x.split("- "),
     markers_used)))
      else:
74
          chr_reg = h5py.File(X_file, 'r')['positions'].attrs['
     chr_regions']
          mk_index= np.array(range(len(markers)),dtype=int)[macs >=
     mac_min]
          chr_pos = np.array([list(map(lambda x: sum(x > chr_reg[:,1]) +
77
      1, mk_index)), markers_used]).T
          my_time = np.repeat((time.time()-start),len(chr_pos))
      pd.DataFrame({
           'chr' : chr_pos[:,0] ,
          'pos' : chr_pos[:,1] ,
81
          'pval': output[:,0] ,
          'mac' : np.array(macs[macs >= mac_min],dtype=np.int) ,
          'eff_size': output[:,1] ,
          'SE' : output[:,2]}).to_csv(out_file,index=False)
  elif perm > 1:
      min_pval = []
      perm_seeds = []
      my\_time = []
      for i in range(perm):
          start_perm = time.time()
          print("Running permutation ", i+1, " of ",perm)
          my_seed = np.asscalar(np.random.randint(9999,size=1))
          perm_seeds.append(my_seed)
          np.random.seed(my_seed)
          Y_perm = np.random.permutation(Y_)
          output = main.gwas(X,K,Y_perm,batch_size)
          min_pval.append(np.min(output[:,0]))
          print("Elapsed time for permuatation", i+1 , " with p_min",
     min_pval[i]," is",": ", round(time.time() - start_perm,2))
          my_time.append(time.time()-start_perm)
      pd.DataFrame({
101
          'time': my_time ,
           'seed': perm_seeds ,
103
           'min_p': min_pval }).to_csv(out_file,index=False)
104
```

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```
print("done")

print("done")

end = time.time()

eltime = np.round(end -start,2)

if eltime <= 59:
    print("Total time elapsed", eltime, "seconds")

elif eltime > 59 and eltime <= 3600:
    print("Total time elapsed", np.round(eltime / 60,2) , "minutes")

elif eltime > 3600 :
    print("Total time elapsed", np.round(eltime / 60 / 60,2), "hours"
    )

print("Total time elapsed", np.round(eltime / 60 / 60,2), "hours"
    )
```

## A.3 main.py

```
import pandas as pd
      import numpy as np
      from scipy.stats import f
      import tensorflow as tf
      import limix
     import herit
      import h5py
      import limix
      import multiprocessing as mlt
10
     def load_and_prepare_data(X_file,Y_file,K_file,m):
11
      type_K = K_file.split(".")[-1]
      type_X = X_file.split(".")[-1]
13
14
      ## load and preprocess genotype matrix
     Y = pd.read_csv(Y_file,engine='python').sort_values(['accession_id
     ']).groupby('accession_id').mean()
     Y = pd.DataFrame({'accession_id': Y.index, 'phenotype_value': Y
     [m]})
     if type_X == 'hdf5' or type_X == 'h5py' :
```

```
SNP = h5py.File(X_file,'r')
          markers= np.asarray(SNP['positions'])
          acc_X = np.asarray(SNP['accessions'][:],dtype=np.int)
      elif type_X == 'csv' :
          X = pd.read_csv(X_file,index_col=0)
          markers = X.columns.values
          acc_X = X.index
          X = np.asarray(X,dtype=np.float32)/2
      else :
          sys.exit("Only hdf5, h5py and csv files are supported")
      if type_K == 'hdf5' or type_K == 'h5py':
          k = h5py.File(K_file,'r')
          acc_K = np.asarray(k['accessions'][:],dtype=np.int)
      elif type_K == 'csv':
33
          k = pd.read_csv(K_file,index_col=0)
          acc_K = k.index
          k = np.array(k, dtype=np.float32)
      acc_Y = np.asarray(Y[['accession_id']]).flatten()
      acc_isec = [isec for isec in acc_X if isec in acc_Y]
      idx_acc = list(map(lambda x: x in acc_isec, acc_X))
      idy_acc = list(map(lambda x: x in acc_isec, acc_Y))
42
      idk_acc = list(map(lambda x: x in acc_isec, acc_K))
43
     Y_ = np.asarray(Y.drop('accession_id',1),dtype=np.float32)[idy_acc
     ,:]
     if type_X == 'hdf5' or type_X == 'h5py' :
47
          X = np.asarray(SNP['snps'][0:(len(SNP['snps'])+1),],dtype=np.
     float32)[:,idx_acc].T
          X = X[np.argsort(acc_X[idx_acc]),:]
          k1 = np.asarray(k['kinship'][:])[idk_acc,:]
          K = k1[:,idk_acc]
          K = K[np.argsort(acc_X[idx_acc]),:]
          K = K[:,np.argsort(acc_X[idx_acc])]
53
      else:
```

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```
X = X[idx_acc,:]
          k1 = k[idk_acc,:]
          K = k1[:,idk_acc]
59
      print("data has been imported")
60
      return X,K,Y_,markers
 def mac_filter(mac_min, X, markers):
      ac1 = np.sum(X,axis=0)
65
      ac0 = X.shape[0] - ac1
      macs = np.minimum(ac1,ac0)
      markers_used = markers[macs >= mac_min]
68
      X = X[:,macs >= mac_min]
69
      return markers_used, X, macs
72 def gwas(X,K,Y,batch_size):
      n_marker = X.shape[1]
73
      n = len(Y)
      ## REML
      K_{stand} = (n-1)/np.sum((np.identity(n) - np.ones((n,n))/n) * K) *
      vg, delta, ve = herit.estimate(Y, "normal", K_stand, verbose = False
77
      print(" Pseudo-heritability is " , vg / (ve + vg + delta))
      print(" Performing GWAS on ", n , " phenotypes and ", n_marker ,"
     markers")
      ## Transform kinship-matrix, phenotypes and estimate intercpt
      Xo = np.ones(K.shape[0]).flatten()
81
      M = np.transpose(np.linalg.inv(np.linalg.cholesky(vg * K_stand +
     ve * np.identity(n))).astype(np.float32)
      Y_t = np.sum(np.multiply(np.transpose(M),Y),axis=1).astype(np.transpose(M),Y)
     float32)
      int_t = np.sum(np.multiply(np.transpose(M),np.ones(n)),axis=1).
     astype(np.float32)
     ## EMMAX Scan
85
```

```
RSS_env = (np.linalg.lstsq(np.reshape(int_t,(n,-1)) , np.reshape(
     Y_t,(n,-1)))[1]).astype(np.float32)
      ## calculate betas and se of betas
      def stderr(a,M,Y_t2d,int_t):
           x = tf.stack((int_t,tf.squeeze(tf.matmul(M.T,tf.reshape(a,(n
      ,-1))))),axis=1)
            coeff = tf.matmul(tf.matmul(tf.linalg.inv(tf.matmul(tf.
      transpose(x),x)),tf.transpose(x)),Y_t2d)
            SSE = tf.reduce_sum(tf.math.square(tf.math.subtract(Y_t,tf.
91
     math.add(tf.math.multiply(x[:,1],coeff[0,0]),tf.math.multiply(x
      [:,1],coeff[1,0])))))
           SE = tf.math.sqrt(SSE/(471-(1+2)))
            StdERR = tf.sqrt(tf.linalg.diag_part(tf.math.multiply(SE , tf
      .linalg.inv(tf.matmul(tf.transpose(x),x))))[1]
            return tf.stack((coeff[1,0],StdERR))
94
      ## calculate residual sum squares
      def rss(a,M,y,int_t):
           x_t = tf.reduce_sum(tf.math.multiply(M.T,a),axis=1)
            lm_res = tf.linalg.lstsq(tf.transpose(tf.stack((int_t,x_t),
      axis=0)), Y_t2d)
           lm_x = tf.concat((tf.squeeze(lm_res),x_t),axis=0)
           return tf.reduce_sum(tf.math.square(tf.math.subtract(tf.
100
      squeeze(Y_t2d),tf.math.add(tf.math.multiply(lm_x[1],lm_x[2:]), tf.
      multiply(lm_x[0],int_t))))
      ## loop over the batches
101
      for i in range(int(np.ceil(n_marker/batch_size))):
          tf.reset_default_graph()
          if n_marker < batch_size:</pre>
104
               X_sub = X
          else:
106
               lower_limit = batch_size * i
               upper_limit = batch_size * i + batch_size
108
               if upper_limit <= n_marker :</pre>
                   X_sub = X[:,lower_limit:upper_limit]
                   print("Working on markers ", lower_limit , " to ",
      upper_limit, " of ", n_marker )
               else:
112
                   X_sub = X[:,lower_limit:]
113
```

A.4. herit.py 85

```
print("Working on markers ", lower_limit , " to ",
114
     n_marker, " of ", n_marker )
          config = tf.ConfigProto()
          n_cores = mlt.cpu_count()
          config.intra_op_parallelism_threads = n_cores
117
          config.inter_op_parallelism_threads = n_cores
118
          sess = tf.Session(config=config)
119
          Y_t2d = tf.cast(tf.reshape(Y_t,(n,-1)),dtype=tf.float32)
          y_tensor = tf.convert_to_tensor(Y_t,dtype = tf.float32)
          StdERR = tf.map_fn(lambda a : stderr(a,M,Y_t2d,int_t), X_sub.T
     )
          R1_full = tf.map_fn(lambda a: rss(a,M,Y_t2d,int_t), X_sub.T)
123
          F_1 = tf.divide(tf.subtract(RSS_env, R1_full),tf.divide(
     R1_full,(n-3))
          if i == 0 :
               output = sess.run(tf.concat([tf.reshape(F_1,(X_sub.shape
      [1],-1)),StdERR],axis=1))
          else :
               tmp = sess.run(tf.concat([tf.reshape(F_1,(X_sub.shape
128
      [1],-1)),StdERR],axis=1))
               output = np.append(output,tmp,axis=0)
129
          sess.close()
130
          F_dist = output[:,0]
      pval = 1 - f.cdf(F_dist,1,n-3)
      output[:,0] = pval
133
      return output
134
136
```

## A.4 herit.py

```
def estimate(y, lik, K, M=None, verbose=True):
from numpy_sugar.linalg import economic_qs
from numpy import pi, var, diag
from glimix_core.glmm import GLMMExpFam
from glimix_core.lmm import LMM
```

```
from limix._data._assert import assert_likelihood
      from limix._data import normalize_likelihood, conform_dataset
      from limix.qtl._assert import assert_finite
      from limix._display import session_block, session_line
10
     lik = normalize_likelihood(lik)
     lik_name = lik[0]
      with session_block("Heritability analysis", disable=not verbose):
          with session_line("Normalising input...", disable=not verbose)
              data = conform_dataset(y, M=M, K=K)
15
          y = data["y"]
          M = data["M"]
          K = data["K"]
          assert_finite(y, M, K)
         if K is not None:
             # K = K / diag(K).mean()
              QS = economic_qs(K)
          else:
              QS = None
          if lik_name == "normal":
              method = LMM(y.values, M.values, QS, restricted=True)
              method.fit(verbose=verbose)
          else:
              method = GLMMExpFam(y, lik, M.values, QS, n_int=500)
              method.fit(verbose=verbose, factr=1e6, pgtol=1e-3)
          g = method.scale * (1 - method.delta)
          e = method.scale * method.delta
          if lik_name == "bernoulli":
              e += pi * pi / 3
          v = var(method.mean())
35
          return g , v , e
```

- Abadi, Martín et al. (2015). TensorFlow: Large-Scale Machine Learning on Heterogeneous Systems. Software available from tensorflow.org. URL: https://www.tensorflow.org/.
- Abadi, Martín et al. (2016). "TensorFlow: A System for Large-scale Machine Learning". In: *Proceedings of the 12th USENIX Conference on Operating Systems Design and Implementation*. OSDI'16. Savannah, GA, USA: USENIX Association, pp. 265–283. ISBN: 978-1-931971-33-1. URL: http://dl.acm.org/citation.cfm?id=3026877.3026899.
- *Albrecht*, Theresa et al. (2011). "Genome-based prediction of testcross values in maize". In: *Theoretical and Applied Genetics* 123.2, p. 339.
- *Allier*, Antoine et al. (2019). "Usefulness Criterion and post-selection Parental Contributions in Multi-parental Crosses: Application to Polygenic Trait Introgression". In: *G3: Genes, Genomes, Genetics* 9.5, pp. 1469–1479.
- Almeida Filho, Janeo Eustáquio de et al. (2019). "Genomic Prediction of Additive and Non-additive Effects Using Genetic Markers and Pedigrees". In: *G3: Genes, Genomes, Genetics* 9.8, pp. 2739–2748. DOI: 10.1534/g3.119.201004. URL: https://doi.org/10.1534.
- Alonso-Blanco, Carlos et al. (2016). "1,135 genomes reveal the global pattern of polymorphism in Arabidopsis thaliana". In: *Cell* 166.2, pp. 481–491.
- Amin, Najaf, Cornelia M Van Duijn, and Yurii S Aulchenko (2007). "A genomic background based method for association analysis in related individuals". In: *PloS one* 2.12, e1274.
- Angermueller, Christof et al. (2016). "Deep learning for computational biology". In: *Molecular systems biology* 12.7, p. 878.
- Ankenbrand, Markus et al. (Jan. 2018). "chloroExtractor: extraction and assembly of the chloroplast genome from whole genome shotgun data". In: *The Journal of*

```
Open Source Software 3.21, p. 464. ISSN: 2475-9066. DOI: 10.21105/joss.00464. URL: http://joss.theoj.org/papers/10.21105/joss.00464.
```

- Ankenbrand, Markus J. and Frank Förster (Apr. 2019). Simulated Arabidopsis thaliana sequencing datasets for chloroplast assembler benchmarking. DOI: 10.5281/zenodo. 2622875. URL: https://doi.org/10.5281/zenodo.2622875.
- Ankenbrand, Markus J. et al. (June 2017). "AliTV—interactive visualization of whole genome comparisons". In: PeerJ Computer Science 3, e116. ISSN: 2376-5992. DOI: 10.7717/peerj-cs.116. URL: https://doi.org/10.7717/peerj-cs.116.
- *Annicchiarico*, Paolo et al. (2015). "Accuracy of genomic selection for alfalfa biomass yield in different reference populations". In: *BMC genomics* 16.1, p. 1020.
- Archibald, John M (2015). "Endosymbiosis and eukaryotic cell evolution". In: *Current Biology* 25.19, R911–R921.
- Auinger, Hans-Jürgen et al. (2016). "Model training across multiple breeding cycles significantly improves genomic prediction accuracy in rye (Secale cereale L.)" In: *Theoretical and Applied Genetics* 129.11, pp. 2043–2053.
- *Azodi*, Christina B et al. (2019). "Benchmarking algorithms for genomic prediction of complex traits". In: *bioRxiv*, p. 614479.
- Bakker, Freek T. et al. (Jan. 2016). "Herbarium genomics: plastome sequence assembly from a range of herbarium specimens using an Iterative Organelle Genome Assembly pipeline". en. In: *Biological Journal of the Linnean Society* 117.1, pp. 33–43. ISSN: 00244066. DOI: 10.1111/bij.12642. URL: https://academic.oup.com/biolinnean/article-lookup/doi/10.1111/bij.12642.
- Belamkar, Vikas et al. (2018). "Genomic Selection in Preliminary Yield Trials in a Winter Wheat Breeding Program". In: *G3: Genes, Genomes, Genetics* 8.8, pp. 2735–2747.

  DOI: 10.1534/g3.118.200415. URL: https://doi.org/10.1534.
- Bendich, Arnold J. (1987). "Why do chloroplasts and mitochondria contain so many copies of their genome?" In: BioEssays 6.6, pp. 279–282. DOI: 10.1002/bies. 950060608. eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1002/bies.950060608. URL: https://onlinelibrary.wiley.com/doi/abs/10.1002/bies.950060608.
- Berardini, Tanya Z. et al. (2015). "The Arabidopsis information resource: Making and mining the "gold standard" annotated reference plant genome". In: genesis 53.8,

```
pp. 474-485. DOI: 10.1002/dvg.22877. eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1002/dvg.22877. URL: https://onlinelibrary.wiley.com/doi/abs/10.1002/dvg.22877.
```

- *Bergstra*, James S et al. (2011). "Algorithms for hyper-parameter optimization". In: *Advances in neural information processing systems*, pp. 2546–2554.
- *Bernal-Vasquez*, Angela-Maria et al. (2014). "The importance of phenotypic data analysis for genomic prediction-a case study comparing different spatial models in rye". In: *BMC genomics* 15.1, p. 646.
- *Bernal-Vasquez*, Angela-Maria et al. (2017). "Genomic prediction in early selection stages using multi-year data in a hybrid rye breeding program". In: *BMC genetics* 18.1, p. 51.
- Bernardo, R (2010). Breeding for quantitative traits in plants. Tech. rep. Stemma Press.
- *Bernardo*, Rex and Jianming *Yu* (2007). "Prospects for genomewide selection for quantitative traits in maize". In: *Crop Science* 47.3, pp. 1082–1090.
- *Biazzi*, Elisa et al. (2017). "Genome-wide association mapping and genomic selection for alfalfa (Medicago sativa) forage quality traits". In: *PLoS One* 12.1, e0169234.
- *Biscarini*, Filippo et al. (2014). "Genome-enabled predictions for binomial traits in sugar beet populations". In: *BMC genetics* 15.1, p. 87.
- *Bock*, Ralph (2017). "Witnessing genome evolution: experimental reconstruction of endosymbiotic and horizontal gene transfer". In: *Annual review of genetics* 51, pp. 1–22.
- Bottou, Léon (1991). "Stochastic gradient learning in neural networks". In: *Proceedings of Neuro-Nimes* 91.8, p. 12.
- Bottou, Léon and Olivier Bousquet (2008). "The Tradeoffs of Large Scale Learning". In: Advances in Neural Information Processing Systems 20 (NIPS 2007). Ed. by J.C. Platt et al. NIPS Foundation (http://books.nips.cc), pp. 161–168. URL: http://leon.bottou.org/papers/bottou-bousquet-2008.
- *Boyle,* Evan A, Yang I *Li*, and Jonathan K *Pritchard* (2017). "An expanded view of complex traits: from polygenic to omnigenic". In: *Cell* 169.7, pp. 1177–1186.
- *Brauner*, Pedro C et al. (2018). "Genomic prediction within and among doubled-haploid libraries from maize landraces". In: *Genetics* 210.4, pp. 1185–1196.
- Brooker, Robert J (1999). Genetics: analysis & principles. Addison-Wesley Reading, MA.

*Bustos-Korts*, Daniela et al. (2016a). "Improvement of predictive ability by uniform coverage of the target genetic space". In: *G3: Genes, Genomes, Genetics* 6.11, pp. 3733–3747.

- (2016b). "Improvement of Predictive Ability by Uniform Coverage of the Target Genetic Space". In: *G3: Genes, Genomes, Genetics* 6.11, pp. 3733–3747. DOI: 10. 1534/g3.116.035410. URL: https://doi.org/10.1534.
- Campos, Gustavo De los and Paulino Perez Rodriguez (2016). BGLR: Bayesian Generalized Linear Regression. R package version 1.0.5. URL: https://CRAN.R-project.org/package=BGLR.
- Chan, Cheong Xin and Mark A. Ragan (2013). "Next-generation phylogenomics". In: Biology direct 8, pp. 3–3. ISSN: 1745-6150. DOI: 10.1186/1745-6150-8-3. URL: https://www.ncbi.nlm.nih.gov/pubmed/23339707.
- Chat, J. et al. (July 2002). "A Case of Chloroplast Heteroplasmy in Kiwifruit (Actinidia deliciosa) That Is Not Transmitted During Sexual Reproduction". In: *Journal of Heredity* 93.4, pp. 293–300. ISSN: 0022-1503. DOI: 10.1093/jhered/93.4.293. eprint: http://oup.prod.sis.lan/jhered/article-pdf/93/4/293/6454216/293.pdf. URL: https://doi.org/10.1093/jhered/93.4.293.
- *Che*, Ronglin et al. (2014). "An adaptive permutation approach for genome-wide association study: evaluation and recommendations for use". In: *BioData mining* 7.1, p. 9.
- Chollet, François et al. (2015). Keras. https://keras.io.
- Coissac, Eric et al. (2016). "From barcodes to genomes: extending the concept of DNA barcoding". In: Molecular Ecology 25.7, pp. 1423–1428. ISSN: 1365-294X. DOI: 10. 1111/mec.13549. URL: https://onlinelibrary.wiley.com/doi/abs/10.1111/mec.13549 (visited on 05/16/2019).
- Collette, Andrew (2013). Python and HDF5. O'Reilly.
- Corriveau, Joseph L. and Annette W. Coleman (1988). "Rapid Screening Method to Detect Potential Biparental Inheritance of Plastid DNA and Results for Over 200 Angiosperm Species". In: American Journal of Botany 75.10, pp. 1443–1458. ISSN: 00029122, 15372197. URL: http://www.jstor.org/stable/2444695.
- *Crossa*, José et al. (2010). "Prediction of genetic values of quantitative traits in plant breeding using pedigree and molecular markers". In: *Genetics*.

Crossa, José et al. (2016). "Genomic Prediction of Gene Bank Wheat Landraces". In: G3: Genes, Genomes, Genetics 6.7, pp. 1819–1834. DOI: 10.1534/g3.116.029637. URL: https://doi.org/10.1534.

- *Crossa*, José et al. (2017). "Genomic selection in plant breeding: Methods, models, and perspectives". In: *Trends in plant science*.
- Cuevas, Jaime et al. (2019a). "Deep Kernel for Genomic and Near Infrared Predictions in Multi-environment Breeding Trials". In: G3: Genes, Genomes, Genetics 9.9, pp. 2913–2924. DOI: 10.1534/g3.119.400493. URL: https://doi.org/10.1534.
- *Cuevas*, Jaime et al. (2019b). "Deep Kernel for Genomic and Near Infrared Predictions in Multi-environment Breeding Trials". In: *G3: Genes, Genomes, Genetics* 9.9, pp. 2913–2924.
- Daniell, Henry et al. (June 23, 2016). "Chloroplast genomes: diversity, evolution, and applications in genetic engineering". In: *Genome Biology* 17. ISSN: 1474-7596. DOI: 10.1186/s13059-016-1004-2. URL: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4918201/ (visited on 05/20/2019).
- Darwin, Charles (1859). On the Origin of Species by Means of Natural Selection. or the Preservation of Favored Races in the Struggle for Life. London: Murray.
- De Los Campos, Gustavo et al. (2009). "Predicting quantitative traits with regression models for dense molecular markers and pedigree". In: *Genetics* 182.1, pp. 375–385.
- *De Rubeis*, Silvia et al. (2014). "Synaptic, transcriptional and chromatin genes disrupted in autism". In: *Nature* 515.7526, p. 209.
- Deiner, Kristy et al. (2017). "Environmental DNA metabarcoding: Transforming how we survey animal and plant communities". In: Molecular Ecology 26.21, pp. 5872–5895. ISSN: 1365-294X. DOI: 10.1111/mec.14350. URL: https://onlinelibrary.wiley.com/doi/abs/10.1111/mec.14350 (visited on 05/21/2019).
- Desta, Zeratsion Abera and Rodomiro Ortiz (2014). "Genomic selection: genomewide prediction in plant improvement". In: Trends in plant science 19.9, pp. 592–601.
- *Dierckxsens*, Nicolas, Patrick *Mardulyn*, and Guillaume *Smits* (Feb. 28, 2017). "NOVO-Plasty: de novo assembly of organelle genomes from whole genome data". In: *Nucleic Acids Research* 45.4, e18–e18. ISSN: 0305-1048. DOI: 10.1093/nar/gkw955.

URL: https://academic.oup.com/nar/article/45/4/e18/2290925 (visited on 01/26/2018).

- Docker Hub Group for Benchmark Project. URL: https://cloud.docker.com/u/chloroextractorteam/.
- Dos Santos, Cicero and Maira Gatti (2014). "Deep convolutional neural networks for sentiment analysis of short texts". In: *Proceedings of COLING 2014, the 25th International Conference on Computational Linguistics: Technical Papers*, pp. 69–78.
- Dozat, Timothy (2016). "Incorporating nesterov momentum into adam". In:
- *El-Dien*, Omnia Gamal et al. (2016). "Implementation of the Realized Genomic Relationship Matrix to Open-Pollinated White Spruce Family Testing for Disentangling Additive from Nonadditive Genetic Effects". In: *G3: Genes, Genomes, Genetics* 6.3, pp. 743–753. DOI: 10.1534/g3.115.025957. URL: https://doi.org/10.1534.
- *Elias*, Ani A et al. (2018a). "Improving genomic prediction in cassava field experiments by accounting for interplot competition". In: *G3: Genes, Genomes, Genetics* 8.3, pp. 933–944.
- (2018b). "Improving genomic prediction in cassava field experiments using spatial analysis". In: *G3: Genes, Genomes, Genetics* 8.1, pp. 53–62.
- Enciso-Rodriguez, Felix et al. (2018). "Genomic selection for late blight and common scab resistance in tetraploid potato (Solanum tuberosum)". In: *G3: Genes, Genomes, Genetics* 8.7, pp. 2471–2481.
- Endelman, Jeffrey B. et al. (Mar. 2018). "Genetic Variance Partitioning and Genome-Wide Prediction with Allele Dosage Information in Autotetraploid Potato". In: Genetics 209.1, pp. 77–87. DOI: 10.1534/genetics.118.300685. URL: https://doi.org/10.1534/genetics.118.300685.
- Falconer, DS and TFC Mackay (1996). "Introduction to quantitative genetics. 1996". In: Harlow, Essex, UK: Longmans Green 3.
- Fan, Jianqing, Fang Han, and Han Liu (2014). "Challenges of big data analysis". In: National science review 1.2, pp. 293–314.
- Feldman, Moshe and Avraham A Levy (2012). "Genome evolution due to allopolyploidization in wheat". In: *Genetics* 192.3, pp. 763–774.

Fisher, Ronald A (1919). "XV.—The correlation between relatives on the supposition of Mendelian inheritance." In: Earth and Environmental Science Transactions of the Royal Society of Edinburgh 52.2, pp. 399–433.

- Förster, Frank and Markus J. Ankenbrand (May 2019). chloroExtractorTeam/benchmark:

  Benchmark container setup v2.0.1. DOI: 10.5281/zenodo.2628061. URL: https://doi.org/10.5281/zenodo.2628061.
- Freudenthal, Jan A. et al. (2019a). "GWAS-Flow: A GPU accelerated framework for efficient permutation based genome-wide association studies". In: DOI: 10.1101/783100. URL: https://doi.org/10.1101.
- *Freudenthal*, Jan A et al. (2019b). "The landscape of chloroplast genome assembly tools". In: *bioRxiv*, p. 665869.
- *Friebe*, B et al. (2000). "Development of a complete set of Triticum aestivum-Aegilops speltoides chromosome addition lines". In: *Theoretical and Applied Genetics* 101.1-2, pp. 51–58.
- *Fuentes*, Ignacia et al. (2014). "Horizontal genome transfer as an asexual path to the formation of new species". In: *Nature* 511.7508, p. 232.
- Gapare, Washington et al. (2018). "Historical Datasets Support Genomic Selection Models for the Prediction of Cotton Fiber Quality Phenotypes Across Multiple Environments". In: G3: Genes, Genomes, Genetics 8.5, pp. 1721–1732.
- *Ghosh,* Sreya et al. (2018). "Speed breeding in growth chambers and glasshouses for crop breeding and model plant research". In: *Nature protocols* 13.12, p. 2944.
- *Gianola*, Daniel (2013). "Priors in whole-genome regression: the Bayesian alphabet returns". In: *Genetics* 194.3, pp. 573–596.
- *Gianola*, Daniel and Johannes BCHM van *Kaam* (2008). "Reproducing kernel Hilbert spaces regression methods for genomic assisted prediction of quantitative traits". In: *Genetics* 178.4, pp. 2289–2303.
- *Gianola*, Daniel and Guilherme JM *Rosa* (2015). "One hundred years of statistical developments in animal breeding". In: *Annu. Rev. Anim. Biosci.* 3.1, pp. 19–56.
- Gianola, Daniel et al. (2009). "Additive genetic variability and the Bayesian alphabet". In: Genetics 183.1, pp. 347–363.
- Gianola, Daniel et al. (2016). "Genome-Wide Association Studies with a Genomic Relationship Matrix: A Case Study with Wheat and Arabidopsis". In: G3: Genes,

```
Genomes, Genetics 6.10, pp. 3241–3256. DOI: 10.1534/g3.116.034256. URL: https://doi.org/10.1534.
```

- GitHub Repository for Benchmark Project. URL: https://github.com/chloroExtractorTeam/benchmark.
- Glorot, Xavier, Antoine Bordes, and Yoshua Bengio (2011). "Deep sparse rectifier neural networks". In: *Proceedings of the fourteenth international conference on artificial intelligence and statistics*, pp. 315–323.
- Goddard, Michael E and Ben J Hayes (2009). "Mapping genes for complex traits in domestic animals and their use in breeding programmes". In: *Nature Reviews Genetics* 10.6, p. 381.
- Goddard, Michael E, Ben J Hayes, and Theo HE Meuwissen (2011). "Using the genomic relationship matrix to predict the accuracy of genomic selection". In: Journal of animal breeding and genetics 128.6, pp. 409–421.
- González-Camacho, JM et al. (2012). "Genome-enabled prediction of genetic values using radial basis function neural networks". In: *Theoretical and Applied Genetics* 125.4, pp. 759–771.
- *González-Camacho*, Juan Manuel et al. (2016). "Genome-enabled prediction using probabilistic neural network classifiers". In: *BMC genomics* 17.1, p. 208.
- González-Camacho, Juan Manuel et al. (2018). "Applications of machine learning methods to genomic selection in breeding wheat for rust resistance". In: *The plant genome* 11.2.
- Goodfellow, Ian, Yoshua Bengio, and Aaron Courville (2016). Deep learning. MIT press.
- *Gouy*, Matthieu et al. (2013). "Experimental assessment of the accuracy of genomic selection in sugarcane". In: *Theoretical and applied genetics* 126.10, pp. 2575–2586.
- Green, Beverley R. (2011). "Chloroplast genomes of photosynthetic eukaryotes". In: The Plant Journal 66.1, pp. 34–44. ISSN: 1365-313X. DOI: 10.1111/j.1365-313X. 2011.04541.x. URL: https://onlinelibrary.wiley.com/doi/abs/10.1111/j. 1365-313X.2011.04541.x (visited on 05/16/2019).
- *Grenier*, Cécile et al. (2015). "Accuracy of genomic selection in a rice synthetic population developed for recurrent selection breeding". In: *PloS one* 10.8, e0136594.

*Grinberg*, Nastasiya F, Oghenejokpeme I *Orhobor*, and Ross D *King* (2018). "An Evaluation of Machine-learning for Predicting Phenotype: Studies in Yeast, Rice and Wheat". In: *BioRxiv*, p. 105528.

- *Guo*, Zhigang et al. (2013). "Accuracy of across-environment genome-wide prediction in maize nested association mapping populations". In: *G3: Genes, Genomes, Genetics* 3.2, pp. 263–272.
- Habier, David, Rohan L Fernando, and Jack CM Dekkers (2007). "The impact of genetic relationship information on genome-assisted breeding values". In: Genetics 177.4, pp. 2389–2397.
- *Habier*, David et al. (2011). "Extension of the Bayesian alphabet for genomic selection". In: *BMC bioinformatics* 12.1, p. 186.
- *Hahnloser*, Richard HR et al. (2000). "Digital selection and analogue amplification coexist in a cortex-inspired silicon circuit". In: *Nature* 405.6789, p. 947.
- Harris, BL, DL Johnson, RJ Spelman, et al. (2009). "Genomic selection in New Zealand and the implications for national genetic evaluation." In: ICAR Technical Series 13, pp. 325–330.
- Hassen, Manel Ben et al. (May 2018). "Genomic Prediction Accounting for Genotype by Environment Interaction Offers an Effective Framework for Breeding Simultaneously for Adaptation to an Abiotic Stress and Performance Under Normal Cropping Conditions in Rice". In: *G3: Genes, Genomes, Genetics* 8.7, pp. 2319–2332. DOI: 10.1534/g3.118.200098. URL: https://doi.org/10.1534/g3.118.200098.
- *Hawkins*, Charles and Long-Xi *Yu* (2018). "Recent progress in alfalfa (Medicago sativa L.) genomics and genomic selection". In: *The Crop Journal* 6.6, pp. 565–575.
- *Hayes*, Ben and Mike *Goddard* (2010). "Genome-wide association and genomic selection in animal breeding". In: *Genome* 53.11, pp. 876–883.
- *Hayes*, BJ, ME *Goddard*, et al. (2001). "Prediction of total genetic value using genomewide dense marker maps". In: *Genetics* 157.4, pp. 1819–1829.
- He, Kaiming et al. (2016). "Deep residual learning for image recognition". In: *Proceedings of the IEEE conference on computer vision and pattern recognition*, pp. 770–778.

Heffner, Elliot L, Jean-Luc Jannink, and Mark E Sorrells (2011). "Genomic selection accuracy using multifamily prediction models in a wheat breeding program". In: *The Plant Genome* 4.1, pp. 65–75.

- Heffner, Elliot L et al. (2010). "Plant breeding with genomic selection: gain per unit time and cost". In: *Crop science* 50.5, pp. 1681–1690.
- *Henderson*, Charles R (1975). "Best linear unbiased estimation and prediction under a selection model". In: *Biometrics*, pp. 423–447.
- *Hinton*, Geoffrey, Nitish *Srivastava*, and Kevin *Swersky* (2012). "Neural networks for machine learning lecture 6a overview of mini-batch gradient descent". In: *Cited on* 14, p. 8.
- Hirschhorn, Joel N. and Mark J. Daly (2005). "Genome-wide association studies for common diseases and complex traits". In: Nature Reviews Genetics 6.2, pp. 95– 108. ISSN: 1471-0064. DOI: 10.1038/nrg1521. URL: https://doi.org/10.1038/ nrg1521.
- Holliday, Jason A., Tongli Wang, and Sally Aitken (2012). "Predicting Adaptive Phenotypes From Multilocus Genotypes in Sitka Spruce (Picea sitchensis) Using Random Forest". In: *G3: Genes, Genomes, Genetics* 2.9, pp. 1085–1093. DOI: 10. 1534/g3.112.002733. URL: https://doi.org/10.1534.
- Howard, Réka et al. (2019). "Joint Use of Genome, Pedigree, and Their Interaction with Environment for Predicting the Performance of Wheat Lines in New Environments". In: *G3: Genes, Genomes, Genetics* 9.9, pp. 2925–2934. DOI: 10.1534/g3. 119.400508. URL: https://doi.org/10.1534.
- *Hu*, Yaodong et al. (2015). "Prediction of plant height in Arabidopsis thaliana using DNA methylation data". In: *Genetics* 201.2, pp. 779–793.
- *Isik*, F (2013). *Genomic Relationships and GBLUP*.
- *Jan*, Habib U et al. (2016). "Genomic prediction of testcross performance in canola (Brassica napus)". In: *PLoS One* 11.1, e0147769.
- *Janocha*, Katarzyna and Wojciech Marian *Czarnecki* (2017). "On loss functions for deep neural networks in classification". In: *arXiv preprint arXiv:1702.05659*.
- *Jaramillo-Correa*, Juan-Pablo et al. (2014). "Molecular Proxies for Climate Maladaptation in a Long-Lived Tree (Pinus pinaster Aiton, Pinaceae)". In: *Genetics* 199.3,

- pp. 793-807. DOI: 10.1534/genetics.114.173252. URL: https://doi.org/10.1534.
- Jarquin, Diego, James Specht, and Aaron Lorenz (2016). "Prospects of Genomic Prediction in the USDA Soybean Germplasm Collection: Historical Data Creates Robust Models for Enhancing Selection of Accessions". In: *G3: Genes, Genomes, Genetics* 6.8, pp. 2329–2341. DOI: 10.1534/g3.116.031443. URL: https://doi.org/10.1534.
- Jette, Morris A., Andy B. Yoo, and Mark Grondona (2002). "SLURM: Simple Linux Utility for Resource Management". In: In Lecture Notes in Computer Science: Proceedings of Job Scheduling Strategies for Parallel Processing (JSSPP) 2003. Springer-Verlag, pp. 44–60.
- *Jiang*, Yong and Jochen C *Reif* (2015). "Modeling epistasis in genomic selection". In: *Genetics* 201.2, pp. 759–768.
- *Jin*, Jian-Jun et al. (Mar. 2018). "GetOrganelle: a simple and fast pipeline for de novo assembly of a complete circular chloroplast genome using genome skimming data". In: *bioRxiv*. DOI: 10.1101/256479. URL: http://biorxiv.org/lookup/doi/10.1101/256479.
- *Kadam*, Dnyaneshwar C et al. (2016). "Genomic prediction of single crosses in the early stages of a maize hybrid breeding pipeline". In: *G3: Genes, Genomes, Genetics* 6.11, pp. 3443–3453.
- *Kainer*, David et al. (2018). "Accuracy of Genomic Prediction for Foliar Terpene Traits in Eucalyptus polybractea". In: *G3: Genes, Genomes, Genetics* 8.8, pp. 2573–2583. DOI: 10.1534/g3.118.200443. URL: https://doi.org/10.1534.
- *Kalo*, P et al. (2004). "Comparative mapping between Medicago sativa and Pisum sativum". In: *Molecular Genetics and Genomics* 272.3, pp. 235–246.
- *Kang*, Hyun Min et al. (2010). "Variance component model to account for sample structure in genome-wide association studies". In: *Nature genetics* 42.4, p. 348.
- *Kärkkäinen*, Hanni P and Mikko J *Sillanpää* (2012). "Back to basics for Bayesian model building in genomic selection". In: *Genetics* 191.3, pp. 969–987.
- *Kim*, Sung et al. (2007). "Recombination and linkage disequilibrium in Arabidopsis thaliana". In: *Nature genetics* 39.9, p. 1151.

*Kingma*, Diederik P and Jimmy *Ba* (2014). "Adam: A method for stochastic optimization". In: *arXiv preprint arXiv:1412.6980*.

- *Kingsolver*, Joel G et al. (2001). "The strength of phenotypic selection in natural populations". In: *The American Naturalist* 157.3, pp. 245–261.
- Koch, Marcus A and Michaela *Matschinger* (2007). "Evolution and genetic differentiation among relatives of Arabidopsis thaliana". In: *Proceedings of the National Academy of Sciences* 104.15, pp. 6272–6277.
- *Korte,* Arthur and Ashley *Farlow* (2013). "The advantages and limitations of trait analysis with GWAS: a review". In: *Plant methods* 9.1, p. 29.
- *Korte,* Arthur et al. (2012). "A mixed-model approach for genome-wide association studies of correlated traits in structured populations". In: *Nature genetics* 44.9, p. 1066.
- Krause, Margaret R. et al. (2019). "Hyperspectral Reflectance-Derived Relationship Matrices for Genomic Prediction of Grain Yield in Wheat". In: *G3: Genes, Genomes, Genetics*, g3.200856.2018. DOI: 10.1534/g3.118.200856. URL: https://doi.org/10.1534.
- Kumar, Rachana A., Delene J. Oldenburg, and Arnold J. Bendich (Sept. 2014). "Changes in DNA damage, molecular integrity, and copy number for plastid DNA and mitochondrial DNA during maize development". In: Journal of Experimental Botany 65.22, pp. 6425–6439. ISSN: 0022-0957. DOI: 10.1093/jxb/eru359. eprint: http://oup.prod.sis.lan/jxb/article-pdf/65/22/6425/16935653/eru359.pdf. URL: https://doi.org/10.1093/jxb/eru359.
- Kumar, Satish et al. (2015). "Genome-Enabled Estimates of Additive and Nonadditive Genetic Variances and Prediction of Apple Phenotypes Across Environments". In: *G3: Genes, Genomes, Genetics* 5.12, pp. 2711–2718. DOI: 10.1534/g3. 115.021105. URL: https://doi.org/10.1534.
- *Kurtzer*, Gregory M, Vanessa *Sochat*, and Michael W *Bauer* (2017). "Singularity: Scientific containers for mobility of compute". In: *PloS one* 12.5, e0177459.
- *Kutschera*, Ulrich and Karl J *Niklas* (2005). "Endosymbiosis, cell evolution, and speciation". In: *Theory in Biosciences* 124.1, pp. 1–24.
- *Lan*, Kun et al. (2018). "A survey of data mining and deep learning in bioinformatics". In: *Journal of medical systems* 42.8, p. 139.

*LeCun*, Yann, Yoshua *Bengio*, and Geoffrey *Hinton* (2015). "Deep learning". In: *nature* 521.7553, p. 436.

- Legarra, Andres, Danila A.L Lourenco, and Zulma G. Vitezica (2018). "Bases for Genomic Prediction". In:
- *Lehermeier*, Christina et al. (2014). "Usefulness of multiparental populations of maize (Zea mays L.) for genome-based prediction". In: *Genetics* 198.1, pp. 3–16.
- Leinonen, Rasko et al. (Nov. 2010). "The Sequence Read Archive". In: Nucleic Acids Research 39.suppl\_1, pp. D19–D21. ISSN: 0305-1048. DOI: 10.1093/nar/gkq1019. eprint: http://oup.prod.sis.lan/nar/article-pdf/39/suppl\\_1/D19/7624335/gkq1019.pdf. URL: https://doi.org/10.1093/nar/gkq1019.
- Leutenegger, Anne-Louise et al. (2003). "Estimation of the inbreeding coefficient through use of genomic data". In: *The American Journal of Human Genetics* 73.3, pp. 516–523.
- *Li*, Bo et al. (2018). "Genomic prediction of breeding values using a subset of SNPs identified by three machine learning methods". In: *Frontiers in genetics* 9, p. 237.
- *Li*, Heng (2018). "Minimap2: pairwise alignment for nucleotide sequences". In: *Bioinformatics* 34.18, pp. 3094–3100.
- *Li*, Jia-Yang, Jun *Wang*, and Robert S *Zeigler* (2014). "The 3,000 rice genomes project: new opportunities and challenges for future rice research". In: *GigaScience* 3.1, p. 8.
- *Li*, Peijin et al. (2014). "Multiple FLC haplotypes defined by independent cis-regulatory variation underpin life history diversity in Arabidopsis thaliana". In: *Genes & Development* 28.15, pp. 1635–1640.
- *Li*, Xuehui and E Charles *Brummer* (2012). "Applied genetics and genomics in alfalfa breeding". In: *Agronomy* 2.1, pp. 40–61.
- *Li*, Xuehui et al. (2015). "Genomic prediction of biomass yield in two selection cycles of a tetraploid alfalfa breeding population". In: *The Plant Genome* 8.2.
- Lippert, Christoph et al. (2014). "LIMIX: genetic analysis of multiple traits". In: bioRxiv. DOI: 10.1101/003905. eprint: https://www.biorxiv.org/content/early/2014/05/22/003905.full.pdf. URL: https://www.biorxiv.org/content/early/2014/05/22/003905.

Litjens, Geert et al. (2017). "A survey on deep learning in medical image analysis". In: Medical image analysis 42, pp. 60–88.

- Lopez-Cruz, Marco et al. (2015). "Increased Prediction Accuracy in Wheat Breeding Trials Using a Marker x Environment Interaction Genomic Selection Model". In: G3: Genes, Genomes, Genetics 5.4, pp. 569–582. DOI: 10.1534/g3.114.016097. URL: https://doi.org/10.1534.
- Luo, Xiang et al. (2017). "Genomic prediction of genotypic effects with epistasis and environment interactions for yield-related traits of rapeseed (Brassica napus L.)" In: Frontiers in genetics 8, p. 15.
- Lynch, Michael, Bruce Walsh, et al. (1998). *Genetics and analysis of quantitative traits*. Vol. 1. Sinauer Sunderland, MA.
- *Ma*, Wenlong et al. (2017). "DeepGS: Predicting phenotypes from genotypes using Deep Learning". In: *bioRxiv*, p. 241414.
- *Mamoshina*, Polina et al. (2016). "Applications of deep learning in biomedicine". In: *Molecular pharmaceutics* 13.5, pp. 1445–1454.
- Martin, William et al. (Sept. 17, 2002). "Evolutionary analysis of Arabidopsis, cyanobacterial, and chloroplast genomes reveals plastid phylogeny and thousands of cyanobacterial genes in the nucleus". In: Proceedings of the National Academy of Sciences of the United States of America 99.19, pp. 12246–12251. ISSN: 0027-8424. DOI: 10.1073/pnas.182432999. URL: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC129430/ (visited on 05/20/2019).
- *Martini*, Johannes WR et al. (2017). "Genomic prediction with epistasis models: on the marker-coding-dependent performance of the extended GBLUP and properties of the categorical epistasis model (CE)". In: *BMC bioinformatics* 18.1, p. 3.
- *Marulanda*, Jose J et al. (2016). "Optimum breeding strategies using genomic selection for hybrid breeding in wheat, maize, rye, barley, rice and triticale". In: *Theoretical and applied genetics* 129.10, pp. 1901–1913.
- Marvin, Minsky and Papert Seymour (1969). Perceptrons.
- McKain, Michael and Afinit (Sept. 2017). Mrmckain/Fast-Plast: Fast-Plast V.1.2.6. DOI: 10.5281/zenodo.973887. URL: https://zenodo.org/record/973887.
- Mckinney, Brett and Nicholas Pajewski (2012). "Six degrees of epistasis: statistical network models for GWAS". In: Frontiers in genetics 2, p. 109.

McKinney, Wes (2010). "Data Structures for Statistical Computing in Python". In: Proceedings of the 9th Python in Science Conference. Ed. by Stéfan van der Walt and Jarrod Millman, pp. 51 –56.

- *Mereschkowsky*, Constantin (1905). "Uber natur und ursprung der chromatophoren im pflanzenreiche". In: *Biologisches Centralblatt* 25, pp. 293–604.
- *Merkel*, Dirk (2014). "Docker: lightweight linux containers for consistent development and deployment". In: *Linux Journal* 2014.239, p. 2.
- Michie, Donald, David J Spiegelhalter, CC Taylor, et al. (1994). "Machine learning". In: Neural and Statistical Classification 13.
- *Min*, Seonwoo, Byunghan *Lee*, and Sungroh *Yoon* (2017). "Deep learning in bioinformatics". In: *Briefings in bioinformatics* 18.5, pp. 851–869.
- *Misztal*, I et al. (2013). "Methods to approximate reliabilities in single-step genomic evaluation". In: *Journal of Dairy Science* 96.1, pp. 647–654.
- *Moeinizade*, Saba et al. (2019). "Optimizing Selection and Mating in Genomic Selection with a Look-Ahead Approach: An Operations Research Framework". In: *G3: Genes, Genomes, Genetics*, g3–200842.
- Momen, Mehdi et al. (Aug. 2019). "Predicting Longitudinal Traits Derived from High-Throughput Phenomics in Contrasting Environments Using Genomic Legendre Polynomials and B-Splines". In: *G3: Genes, Genomes, Genetics* 9.10, pp. 3369–3380. DOI: 10.1534/g3.119.400346. URL: https://doi.org/10.1534/g3.119.400346.
- *Montesinos-López*, Osval A et al. (2015). "Threshold models for genome-enabled prediction of ordinal categorical traits in plant breeding". In: *G3: Genes, Genomes, Genetics* 5.2, pp. 291–300.
- Montesinos-López, Osval A et al. (2019a). "A benchmarking between deep learning, support vector machine and Bayesian threshold best linear unbiased prediction for predicting ordinal traits in plant breeding". In: *G3: Genes, Genomes, Genetics* 9.2, pp. 601–618.
- (2019b). "New deep learning genomic-based prediction model for multiple traits with binary, ordinal, and continuous phenotypes". In: *G3: Genes, Genomes, Genetics* 9.5, pp. 1545–1556.
- *Morota,* Gota and Daniel *Gianola* (2014). "Kernel-based whole-genome prediction of complex traits: a review". In: *Frontiers in genetics* 5, p. 363.

Moser, Gerhard et al. (2009). "A comparison of five methods to predict genomic breeding values of dairy bulls from genome-wide SNP markers". In: *Genetics Selection Evolution* 41.1, p. 56.

- *Mousseau*, Timothy A and Derek A *Roff* (1987). "Natural selection and the heritability of fitness components". In: *Heredity* 59.2, p. 181.
- National Center for Biotechnology Information. NCBI Taxonomy. Accessed: 2019-10-01. URL: https://www.ncbi.nlm.nih.gov/Taxonomy/taxonomyhome.html/.
- *Neyhart*, Jeffrey, Aaron J *Lorenz*, and Kevin P *Smith* (2019). "Multi-Trait Improvement by Predicting Genetic Correlations in Breeding Crosses". In: *bioRxiv*, p. 593210.
- *Nguyen*, Derrick and Bernard *Widrow* (1990). "The truck backer-upper: An example of self-learning in neural networks". In: *Advanced neural computers*. Elsevier, pp. 11–19.
- *Nordborg*, Magnus et al. (2002). "The extent of linkage disequilibrium in Arabidopsis thaliana". In: *Nature genetics* 30.2, p. 190.
- Norman, Adam et al. (2018). "Optimising Genomic Selection in Wheat: Effect of Marker Density, Population Size and Population Structure on Prediction Accuracy". In: *G3: Genes, Genomes, Genetics* 8.9, pp. 2889–2899. DOI: 10.1534/g3.118. 200311. URL: https://doi.org/10.1534.
- Oakey, Helena et al. (2016). "Genomic selection in multi-environment crop trials". In: *G3: Genes, Genomes, Genetics* 6.5, pp. 1313–1326.
- Ogutu, Joseph O, Hans-Peter *Piepho*, and Torben *Schulz-Streeck* (2011). "A comparison of random forests, boosting and support vector machines for genomic selection". In: *BMC proceedings*. Vol. 5. 3. BioMed Central, S11.
- Ohyama, Kanji et al. (Aug. 1986). "Chloroplast gene organization deduced from complete sequence of liverwort Marchantia polymorpha chloroplast DNA". In: *Nature* 322.6079, p. 572. ISSN: 1476-4687. DOI: 10.1038/322572a0. URL: https://www.nature.com/articles/322572a0 (visited on 05/20/2019).
- Olejniczak, Szymon Adam et al. (2016). "Chloroplasts: state of research and practical applications of plastome sequencing". In: *Planta* 244.3, pp. 517–527.
- Oliphant, Travis E (2006). A guide to NumPy. Vol. 1. Trelgol Publishing USA.

Ovenden, Ben et al. (2018). "Accounting for Genotype-by-Environment Interactions and Residual Genetic Variation in Genomic Selection for Water-Soluble Carbohydrate Concentration in Wheat". In: *G3: Genes, Genomes, Genetics* 8.6, pp. 1909–1919. DOI: 10.1534/g3.118.200038. URL: https://doi.org/10.1534.

- *Owens*, Brenda F et al. (2014). "A foundation for provitamin A biofortification of maize: genome-wide association and genomic prediction models of carotenoid levels". In: *Genetics* 198.4, pp. 1699–1716.
- Ozkan, Hakan, Avraham A *Levy*, and Moshe *Feldman* (2001). "Allopolyploidy-induced rapid genome evolution in the wheat (Aegilops–Triticum) group". In: *The Plant Cell* 13.8, pp. 1735–1747.
- Palmer, Jeffrey D. (1985). "COMPARATIVE ORGANIZATION OF CHLOROPLAST GENOMES". In: Annual Review of Genetics 19.1. PMID: 3936406, pp. 325–354. DOI: 10.1146/annurev.ge.19.120185.001545. eprint: https://doi.org/10.1146/annurev.ge.19.120185.001545. URL: https://doi.org/10.1146/annurev.ge.19.120185.001545.
- *Peiffer*, Jason A et al. (2014). "The genetic architecture of maize height". In: *Genetics* 196.4, pp. 1337–1356.
- *Polyak*, Boris T (1964). "Some methods of speeding up the convergence of iteration methods". In: *USSR Computational Mathematics and Mathematical Physics* 4.5, pp. 1–17.
- Poudel, Hari P. et al. (2019). "Genomic Prediction for Winter Survival of Lowland Switchgrass in the Northern USA". In: *G3: Genes, Genomes, Genetics*, g3.400094.2019. DOI: 10.1534/g3.119.400094. URL: https://doi.org/10.1534.
- Purcell, Shaun et al. (2007). "PLINK: a tool set for whole-genome association and population-based linkage analyses". In: *The American journal of human genetics* 81.3, pp. 559–575.
- *Purcell,* Shaun M et al. (2014). "A polygenic burden of rare disruptive mutations in schizophrenia". In: *Nature* 506.7487, p. 185.
- *Qian*, Lunwen, Wei *Qian*, and Rod J *Snowdon* (2014). "Sub-genomic selection patterns as a signature of breeding in the allopolyploid Brassica napus genome". In: *BMC genomics* 15.1, p. 1170.

*Qiu*, Zhixu et al. (2016). "Application of machine learning-based classification to genomic selection and performance improvement". In: *International Conference on Intelligent Computing*. Springer, pp. 412–421.

- R Core Team (2019). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing. Vienna, Austria. URL: https://www.R-project.org/.
- Rampasek, Ladislav and Anna Goldenberg (2016). "Tensorflow: Biology's gateway to deep learning?" In: Cell systems 2.1, pp. 12–14.
- Ramstein, Guillaume P. and Michael D. Casler (2019). "Extensions of BLUP Models for Genomic Prediction in Heterogeneous Populations: Application in a Diverse Switchgrass Sample". In: *G3: Genes, Genomes, Genetics*, g3.200969.2018. DOI: 10. 1534/g3.118.200969. URL: https://doi.org/10.1534.
- Ramstein, Guillaume P. et al. (2016). "Accuracy of Genomic Prediction in Switchgrass (Panicum virgatumL.) Improved by Accounting for Linkage Disequilibrium". In: G3: Genes, Genomes, Genetics 6.4, pp. 1049–1062. DOI: 10.1534/g3.115.024950. URL: https://doi.org/10.1534.
- Ratcliffe, Blaise et al. (2017). "Single-Step BLUP with Varying Genotyping Effort in Open-Pollinated Picea glauca". In: *G3: Genes, Genomes, Genetics* 7.3, pp. 935–942. DOI: 10.1534/g3.116.037895. URL: https://doi.org/10.1534.
- Resende, M. F. R. et al. (2012). "Accuracy of Genomic Selection Methods in a Standard Data Set of Loblolly Pine (Pinus taeda L.)" In: *Genetics* 190.4, pp. 1503–1510. DOI: 10.1534/genetics.111.137026. URL: https://doi.org/10.1534.
- *Riedelsheimer*, Christian et al. (2013). "Genomic predictability of interconnected biparental maize populations". In: *Genetics* 194.2, pp. 493–503.
- *Rincent*, Renaud et al. (2012). "Maximizing the reliability of genomic selection by optimizing the calibration set of reference individuals: comparison of methods in two diverse groups of maize inbreds (Zea mays L.)" In: *Genetics* 192.2, pp. 715–728.
- Rincent, Renaud et al. (2018). "Phenomic Selection Is a Low-Cost and High-Throughput Method Based on Indirect Predictions: Proof of Concept on Wheat and Poplar".

  In: G3: Genes, Genomes, Genetics, g3.200760.2018. DOI: 10.1534/g3.118.200760.

  URL: https://doi.org/10.1534.

Ritchie, Marylyn D and Kristel Van Steen (2018). "The search for gene-gene interactions in genome-wide association studies: challenges in abundance of methods, practical considerations, and biological interpretation". In: Annals of translational medicine 6.8.

- Rocaps Lab. CpBase. Accessed: 2019-04-01, Version: 8/20/2017. URL: http://rocaplab.ocean.washington.edu/old\_website/tools/cpbase.
- Rosenblatt, Frank (1961). Principles of neurodynamics. perceptrons and the theory of brain mechanisms. Tech. rep. Cornell Aeronautical Lab Inc Buffalo NY.
- *Ruder*, Sebastian (2016). "An overview of gradient descent optimization algorithms". In: *arXiv preprint arXiv:1609.04747*.
- Rumelhart, David E, Geoffrey E Hinton, Ronald J Williams, et al. (1988). "Learning representations by back-propagating errors". In: Cognitive modeling 5.3, p. 1.
- Sancho, Rubén et al. (June 2018). "Comparative plastome genomics and phylogenomics of *Brachypodium*: flowering time signatures, introgression and recombination in recently diverged ecotypes". en. In: *New Phytologist* 218.4, pp. 1631–1644. ISSN: 0028646X. DOI: 10.1111/nph.14926. URL: http://doi.wiley.com/10.1111/nph.14926.
- Scarcelli, N. et al. (2016). "Intra-individual polymorphism in chloroplasts from NGS data: where does it come from and how to handle it?" In: Molecular Ecology Resources 16.2, pp. 434–445. DOI: 10.1111/1755-0998.12462. eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1111/1755-0998.12462. URL: https://onlinelibrary.wiley.com/doi/abs/10.1111/1755-0998.12462.
- Schmidhuber, Jürgen (2015). "Deep learning in neural networks: An overview". In: Neural networks 61, pp. 85–117.
- Schopp, Pascal et al. (2017a). "Accuracy of genomic prediction in synthetic populations depending on the number of parents, relatedness, and ancestral linkage disequilibrium". In: *Genetics* 205.1, pp. 441–454.
- *Schopp*, Pascal et al. (2017b). "Genomic prediction within and across biparental families: means and variances of prediction accuracy and usefulness of deterministic equations". In: *G3: Genes, Genomes, Genetics* 7.11, pp. 3571–3586.

Schrag, Tobias A et al. (2018). "Beyond genomic prediction: combining different types of omics data can improve prediction of hybrid performance in maize". In: *Genetics* 208.4, pp. 1373–1385.

- Segura, Vincent et al. (2012). "An efficient multi-locus mixed-model approach for genome-wide association studies in structured populations". In: *Nature genetics* 44.7, p. 825.
- Seren, Ümit et al. (2016). "AraPheno: a public database for Arabidopsis thaliana phenotypes". In: *Nucleic acids research*, gkw986.
- Shen, Wei et al. (Oct. 2016). "SeqKit: A Cross-Platform and Ultrafast Toolkit for FAS-TA/Q File Manipulation". In: PLOS ONE 11.10, pp. 1–10. DOI: 10.1371/journal.pone.0163962. URL: https://doi.org/10.1371/journal.pone.0163962.
- *Shen*, Xia et al. (2013). "A novel generalized ridge regression method for quantitative genetics". In: *Genetics* 193.4, pp. 1255–1268.
- Shinozaki, K. et al. (1986). "The complete nucleotide sequence of the tobacco chloroplast genome: its gene organization and expression". In: *The EMBO Journal* 5.9, pp. 2043–2049. ISSN: 1460-2075. DOI: 10.1002/j.1460-2075.1986.tb04464.x. URL: https://onlinelibrary.wiley.com/doi/abs/10.1002/j.1460-2075.1986.tb04464.x (visited on 05/20/2019).
- Siva, Nayanah (2008). 1000 Genomes project.
- *Snowdon*, Rod J and Federico L *Iniguez Luy* (2012). "Potential to improve oilseed rape and canola breeding in the genomics era". In: *Plant breeding* 131.3, pp. 351–360.
- Soper, HE et al. (1917). "On the distribution of the correlation coefficient in small samples. Appendix II to the papers of" Student" and RA Fisher". In: *Biometrika* 11.4, pp. 328–413.
- Sousa, Massaine Bandeira e et al. (2017). "Genomic-enabled prediction in maize using kernel models with genotype× environment interaction". In: *G3: Genes, Genomes, Genetics* 7.6, pp. 1995–2014.
- Stegemann, Sandra and Ralph *Bock* (2009). "Exchange of genetic material between cells in plant tissue grafts". In: *science* 324.5927, pp. 649–651.
- Stewart-Brown, Benjamin B. et al. (2019). "Genomic Selection for Yield and Seed Composition Traits Within an Applied Soybean Breeding Program". In: G3: Genes,

```
Genomes, Genetics 9.7, pp. 2253–2265. DOI: 10.1534/g3.118.200917. URL: https://doi.org/10.1534.
```

- Storey, John D. and Robert *Tibshirani* (2003). "Statistical significance for genomewide studies". In: *Proceedings of the National Academy of Sciences* 100.16, pp. 9440–9445. ISSN: 0027-8424. DOI: 10.1073/pnas.1530509100. eprint: https://www.pnas.org/content/100/16/9440.full.pdf. URL: https://www.pnas.org/content/100/16/9440.
- Stringer, Sven et al. (2011). "Underestimated effect sizes in GWAS: fundamental limitations of single SNP analysis for dichotomous phenotypes". In: *PloS one* 6.11, e27964.
- Sukumaran, Sivakumar et al. (2016). "Genomic Prediction with Pedigree and Genotype Environment Interaction in Spring Wheat Grown in South and West Asia, North Africa, and Mexico". In: *G3: Genes, Genomes, Genetics* 7.2, pp. 481–495. DOI: 10.1534/g3.116.036251. URL: https://doi.org/10.1534.
- *Tang*, Chunlao et al. (2007). "The evolution of selfing in Arabidopsis thaliana". In: *Science* 317.5841, pp. 1070–1072.
- *Technow*, Frank, Anna *Bürger*, and Albrecht E *Melchinger* (2013). "Genomic prediction of northern corn leaf blight resistance in maize with combined or separated training sets for heterotic groups". In: *G3: Genes, Genomes, Genetics* 3.2, pp. 197–203.
- *Technow*, Frank et al. (2014). "Genome properties and prospects of genomic prediction of hybrid performance in a breeding program of maize". In: *Genetics* 197.4, pp. 1343–1355.
- Tetko, Igor V, David J Livingstone, and Alexander I Luik (1995). "Neural network studies. 1. Comparison of overfitting and overtraining". In: Journal of chemical information and computer sciences 35.5, pp. 826–833.
- Thavamanikumar, Saravanan, Rudy Dolferus, and Bala R. Thumma (2015). "Comparison of Genomic Selection Models to Predict Flowering Time and Spike Grain Number in Two Hexaploid Wheat Doubled Haploid Populations". In: *G3: Genes, Genomes, Genetics* 5.10, pp. 1991–1998. DOI: 10.1534/g3.115.019745. URL: https://doi.org/10.1534.

*Thorwarth*, Patrick, Eltohamy AA *Yousef*, and Karl J *Schmid* (2018). "Genomic Prediction and Association Mapping of Curd-Related Traits in Gene Bank Accessions of Cauliflower". In: *G3: Genes, Genomes, Genetics* 8.2, pp. 707–718.

- *Timpson*, Nicholas J et al. (2018). "Genetic architecture: the shape of the genetic contribution to human traits and disease". In: *Nature Reviews Genetics* 19.2, p. 110.
- Togninalli, Matteo et al. (2017). "The AraGWAS Catalog: a curated and standardized Arabidopsis thaliana GWAS catalog". In: *Nucleic acids research* 46.D1, pp. D1150–D1156.
- Twyford, Alex D. and Rob W. Ness (Sept. 1, 2017). "Strategies for complete plastid genome sequencing". In: Molecular Ecology Resources 17.5, pp. 858–868. ISSN: 1755-0998. DOI: 10.1111/1755-0998.12626. URL: http://onlinelibrary.wiley.com/doi/10.1111/1755-0998.12626/abstract (visited on 01/26/2018).
- *Van Rossum*, Guido and Fred L *Drake Jr* (1995). *Python tutorial*. Centrum voor Wiskunde en Informatica Amsterdam, The Netherlands.
- VanRaden, Paul M (2008). "Efficient methods to compute genomic predictions". In: *Journal of dairy science* 91.11, pp. 4414–4423.
- *VanRaden*, PM et al. (2008). "Reliability of genomic predictions for North American dairy bulls". In: *J. Dairy Sci* 91. Suppl 1, p. 305.
- *Verbyla*, Klara L et al. (2009). "Accuracy of genomic selection using stochastic search variable selection in Australian Holstein Friesian dairy cattle". In: *Genetics research* 91.5, pp. 307–311.
- *Vieira*, IC et al. (2017). "Assessing non-additive effects in GBLUP model". In: *Genetics* and molecular research: GMR 16.2.
- Vinga, Susana et al. (2012). "Pattern matching through Chaos Game Representation: bridging numerical and discrete data structures for biological sequence analysis". In: Algorithms for molecular biology: AMB 7.1. 22551152[pmid], pp. 10–10. ISSN: 1748-7188. DOI: 10.1186/1748-7188-7-10. URL: https://www.ncbi.nlm.nih.gov/pubmed/22551152.
- Walsh, B and M Lynch (2018a). "Short-term Changes in the Mean: 2. Truncation and Threshold Selection". In:
- Walsh, Bruce and Michael Lynch (2018b). Evolution and selection of quantitative traits. Oxford University Press.

Wang, Weiwen et al. (2018). "Assembly of chloroplast genomes with long- and short-read data: a comparison of approaches using Eucalyptus pauciflora as a test case". In: *BMC genomics* 19.1. 30594129[pmid], pp. 977–977. ISSN: 1471-2164. DOI: 10.1186/s12864-018-5348-8. URL: https://www.ncbi.nlm.nih.gov/pubmed/30594129.

- Wang, Yu et al. (2014). "The accuracy of prediction of genomic selection in elite hybrid rye populations surpasses the accuracy of marker-assisted selection and is equally augmented by multiple field evaluation locations and test years". In: *BMC genomics* 15.1, p. 556.
- *Warner*, Brad and Manavendra *Misra* (1996). "Understanding neural networks as statistical tools". In: *The american statistician* 50.4, pp. 284–293.
- *Watson,* Amy et al. (2018). "Speed breeding is a powerful tool to accelerate crop research and breeding". In: *Nature plants* 4.1, p. 23.
- Webb, Sarah (2018). "Deep learning for biology". In: Nature 554.7693.
- Werner, Christian R et al. (2018). "Effective genomic selection in a narrow-genepool crop with low-density markers: Asian rapeseed as an example". In: *The plant genome* 11.2.
- *Wicke*, Susann et al. (July 1, 2011). "The evolution of the plastid chromosome in land plants: gene content, gene order, gene function". In: *Plant Molecular Biology* 76.3, pp. 273–297. ISSN: 1573-5028. DOI: 10.1007/s11103-011-9762-4. URL: https://doi.org/10.1007/s11103-011-9762-4 (visited on 05/16/2019).
- Windhausen, Vanessa S et al. (2012). "Effectiveness of genomic prediction of maize hybrid performance in different breeding populations and environments". In: *G3: Genes, Genomes, Genetics* 2.11, pp. 1427–1436.
- Wright, Sewall (1922). "Coefficients of inbreeding and relationship". In: *The American Naturalist* 56.645, pp. 330–338.
- Würschum, Tobias (2012). "Mapping QTL for agronomic traits in breeding populations". In: *Theoretical and Applied Genetics* 125.2, pp. 201–210.
- Würschum, Tobias, Stefan Abel, and Yusheng Zhao (2014). "Potential of genomic selection in rapeseed (B rassica napus L.) breeding". In: Plant Breeding 133.1, pp. 45–51.

*Würschum,* Tobias et al. (2013). "Genomic selection in sugar beet breeding populations". In: *BMC genetics* 14.1, p. 85.

- *Xavier*, Alencar, William M. *Muir*, and Katy Martin *Rainey* (2016). "Assessing Predictive Properties of Genome-Wide Selection in Soybeans". In: *G3* 6.8, pp. 2611–2616. DOI: 10.1534/g3.116.032268. URL: https://doi.org/10.1534.
- Xiao-Ming, Zheng et al. (May 8, 2017). "Inferring the evolutionary mechanism of the chloroplast genome size by comparing whole-chloroplast genome sequences in seed plants". In: *Scientific Reports* 7.1, p. 1555. ISSN: 2045-2322. DOI: 10.1038/s41598-017-01518-5. URL: https://www.nature.com/articles/s41598-017-01518-5 (visited on 05/16/2019).
- *Xu*, S (2010). "An expectation–maximization algorithm for the Lasso estimation of quantitative trait locus effects". In: *Heredity* 105.5, p. 483.
- Xu, Shizhong (Aug. 2013). "Genetic Mapping and Genomic Selection Using Recombination Breakpoint Data". In: *Genetics* 195.3, pp. 1103–1115. DOI: 10.1534/genetics.113.155309. URL: https://doi.org/10.1534/genetics.113.155309.
- *Yang*, Jian et al. (2010). "Common SNPs explain a large proportion of the heritability for human height". In: *Nature genetics* 42.7, p. 565.
- Zapata-Valenzuela, Jaime et al. (2013). "Genomic Estimated Breeding Values Using Genomic Relationship Matrices in a Cloned Population of Loblolly Pine". In: *G3:* Genes, Genomes, Genetics 3.5, pp. 909–916. DOI: 10.1534/g3.113.005975. URL: https://doi.org/10.1534.
- *Zeiler*, Matthew D (2012). "ADADELTA: an adaptive learning rate method". In: *arXiv* preprint arXiv:1212.5701.
- Zhang, Yan et al. (2018). "Estimation of complex effect-size distributions using summary-level statistics from genome-wide association studies across 32 complex traits".

  In: Nature genetics 50.9, p. 1318.
- *Zhang*, Zhiwu et al. (Mar. 2010). "Mixed linear model approach adapted for genomewide association studies". In: *Nature Genetics* 42.4, pp. 355–360. DOI: 10.1038/ng. 546. URL: https://doi.org/10.1038/ng.546.
- *Zhong*, Shengqiang et al. (2009). "Factors affecting accuracy from genomic selection in populations derived from multiple inbred lines: a barley case study". In: *Genetics* 182.1, pp. 355–364.

*Zhou*, Xiang and Matthew *Stephens* (June 2012). "Genome-wide efficient mixed-model analysis for association studies". In: *Nature Genetics* 44.7, pp. 821–824. DOI: 10.1038/ng.2310. URL: https://doi.org/10.1038/ng.2310.

Žilinskas, A (2006). Practical mathematical optimization: An introduction to basic optimization theory and classical and new gradient-based algorithms.