# QTL mapping in MAGIC populations with R/qtl2

#### Karl Broman

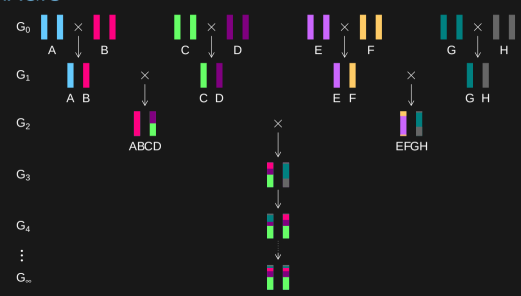
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Slides: kbroman.org/Talk\_MAGIC2021



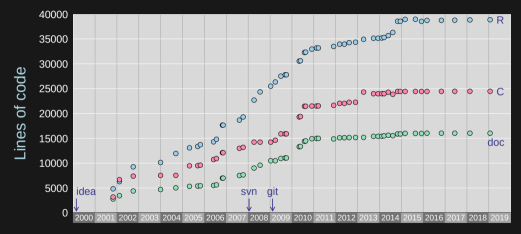
# **MAGIC**



## MAGIC is magic

- Genetic diversity
- ► High-precision mapping
- Predictable linkage disequilibrium
- Phenotype replicates to reduce individual variation
- Pool phenotypes from multiple labs, environments, treatments
- Genotype once
- ▶ Cool name

## 19 years of R/qtl



Year

## R/qtl cross types

- backcross, doubled haploids, haploid
- intercross
- 2-way RIL by selfing or sibling mating
- ▶ phase-known 4-way cross



## R/qtl2 cross types

- backcross, doubled haploids, haploid
- intercross
- ► 2-, 4-, 8-, 16-way RIL by selfing
- ▶ 2-, 4-, 8-way RIL by sibling mating
- ► 2-, 3-, 8-way advanced intercross
- ► 6- and 19-way MAGIC
- Diversity Outbred (DO) mice
- ightharpoonup  $F_1$  of DO imes inbred
- ▶ general RIL or AIL

	А	В	С	D	E	F	G
1	id	bolting_days	seed_weight	seed_area	ttl_seedspfruit	branches	height
2	MAGIC.1	15.33	17.15	0.64	45.11	10.5	NA
3	MAGIC.2	22	22.71	0.75	49.11	4.33	42.33
4	MAGIC.3	23	21.03	0.68	57	4.67	50
5	MAGIC.4	18.67	22.45	0.74	54.33	6.33	NA
6	MAGIC.5	18.67	25.36	0.82	38.33	5.67	42.25
7	MAGIC.6	25	21.53	0.71	52	4.33	NA
8	MAGIC.7	15.33	20.92	0.71	39	4	37.35
9	MAGIC.8	14.33	24.2	0.79	50.56	7.33	43.23
10	MAGIC.9	16.33	18.86	0.63	75.78	5.33	42.4
11	MAGIC.10	30	30.46	0.91	56.33	3	48.5
12	MAGIC.11	14	20.58	0.66	41.56	5.67	36.25
13	MAGIC.12	21.33	19.05	0.67	52.33	4.67	49.35
14	MAGIC.14	18.67	21.14	0.72	49.78	6	48.5
15	MAGIC.15	15	21.71	0.73	49	6.33	41.9
16	MAGIC.16	18.33	21.37	0.71	66.67	3.33	18.17

	Α		В	С	D	Е	F	G	
1	id	t	oolting_days	seed_weight	seed_area	ttl_seedspfruit	branches	height	
2			A	В	С	D	Е	F	G
3	1		marker	MAGIC.1	MAGIC.10	MAGIC.100	MAGIC.101	MAGIC.102	MAGIC.103
4	2		MN1_29291	Α	Α	Α	В	В	Α
5	3		MN1_29716	Α	Α	Α	В	В	Α
6	4		MN1_11290	7 A	Α	Α	В	В	Α
7	5		MASC03771	Α	В	Α	Α	В	Α
8	6		MN1_19778	7 A	В	Α	В	В	Α
9	7		MN1_340810	Α Α	В	Α	Α	Α	Α
10	8		MN1_39510	7 A	Α	Α	Α	В	Α
11	9		MN1_444820	Α	Α	-	Α	Α	Α
12	10		MN1_49420	5 A	Α	Α	Α	Α	Α
13	11		MN1_59286	3 A	В	Α	Α	Α	Α
14	12		MN1_592760	Α Α	Α	Α	Α	Α	Α
15	13		BKN118	Α	Α	Α	Α	Α	Α
16	14		MN1_104242	7 A	В	В	Α	В	В
	15		CRY2_1021	Α	Α	Α	Α	Α	Α
	16		CRY2_429	Α	Α	Α	Α	В	Α

	Α	В	В		С		D		Е				F			G							
1	id	bolting_da	olting_days seed_v		veight	seed_area		ttl_seedspfruit		br	branches		height										
2		А			В		С			D			Е			F			G				
3	1	mar	ker	MA	GIC.	1	MAGIC.10		MAGIC.100		1 (	MAGIC.101			MAGIC.102			MAGIC.103					
4	2		1	Ą	В	С	D	Е	F	G	Н	-1	J	K	L	M	N	0	Р	Q	R	S	Т
5	3	1	ma		Bur	Can	Col	Ct	Edi	Hi	Kn	Ler	Mt	No	Oy	Po	Rsch	Sf	Tsu	Wil	Ws	Wu	Zu
6	4	2	MN1_	29291 29716	В	A	A	В	A	A	В	A	A	A	В	В	A	В	A	A	A	A	A
7	5	4	_	112907	В	A	A	A	В	В	В	В	A	A	В	В	A	В	В	A	A	A	A
8	6	5	MASC	03771	Α	В	Α	Α	В	Α	В	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α
9	7	- 6	_	197787	В	В	Α	Α	Α	В	В	Α	Α	Α	Α	Α	Α	В	Α	Α	Α	В	В
10	8	7 8	_	340810 395107	В	В	A	В	A	B	A B	A	A	A	A	A	A	A	A	A	В	В	A
	9	9	_	144820	A B	A	A	A B	A B	A	A	A	A	A	A	A	A	A	A B	A	A B	A B	A
11	_	10	_	194205	В	A	A	A	В	A	A	A	A	A	A	A	A	A	A	A	A	В	A
12	10	- 11	MN1_5	592863	A	В	Α	В	A	Α	Α	В	Α	Α	Α	Α	Α	Α	В	Α	В	Α	Α
13	11	12	MN1_	592760	В	Α	Α	Α	В	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α
14	12	13	BKN	V118	В	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α
15	13	14	_	042427	Α	В	Α	В	Α	В	В	Α	В	Α	Α	Α	В	Α	Α	В	Α	В	Α
16	14	15	-	_1021	A	A	A	A	В	Α	Α	A	A	A	A	A	A	A	A	A	A	A	A
. 0	15	16	-	2_429	A	A	A B	A	A	B	B A	B A	A	B A	A	A	A B	A	A	A B	B A	A	A
		17		03609	A	A	A	A	A	A	В	B	A	В	В	B	A	A	A	A	B	A	A
	16	19		296068	A	A	A	A	A	A	В	A	A	A	A	A	A	A	A	A	В	В	A
		20	_	399466	Α	A	Α	A	Α	Α	В	Α	Α	A	A	Α	Α	Α	A	-	В	A	Α
		21	AXR'	1_381	Α	В	Α	Α	Α	В	Α	Α	Α	В	Α	Α	Α	Α	Α	Α	В	Α	В
		22	MASC	07424	Α	В	В	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	В	-	Α	Α	Α	Α

	A	В		С		D			Е		F		G									
1	id	bolting_da	vs se	seed weight		seed area		ttl	ttl seedspfruit		branches		height									
2	IQ	A	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	В			C C		D		E		F				G					
3	4							_			+			-					100			
	1	mark		MAGIC			MAGIC.10		MAGIC.100		_	MAGIC.101					MAGIC.103					
4	2		A	В	С	_	D E	F	G	Н	- 1	J	K	L	M	N	0	Р	Q	R	S	Т
5	3	1 2	mark MN1_2		Ca	n (	Col Ct	Fdi	Hi	Kn B	Ler	Mt	No.	Ov	Po B	Rsch	Sf B	Tsu	Wil	Ws	Wu	Zu
6	4	3	MN1_2		1	-						pos			В	A	В	A	A	A	A	A
7	5	4	MN1_11	2		marker			chr						В	A	В	В	A	A	A	A
8	6	5	MASC0	3	2		MN1_29		1	0.029291				A	A	A	A	A	A	A	A	
		- 6	MN1_19	7	3		MN1_29716			1	0.029757			7 A		Α	В	Α	Α	Α	В	В
9	7	7	MN1_34	40 4			MN1_11	2907		1 0.11290			7	Α	Α	Α	Α	Α	В	В	Α	
10	8	8	MN1_39	15	5		MASC03771			1		0.174605		5	Α	Α	Α	Α	Α	Α	Α	Α
11	9	9	MN1_44	14		MN1 19778				1		0.197787		7	Α	Α	Α	В	Α	В	В	Α
12	10	10	MN1_49												Α	Α	Α	Α	Α	Α	В	Α
13	11	11	MN1_59		7	MN1_340				1			0.34081		A	A	A	В	A	В	A	A
		12	MN1_59 BKN1		3		MN1_395			1		0.395107		7	A	A	A	A	A	A	A	A
14	12	14	MN1 10		9	MN1_444		4820	20 1		0.444764		4	A	В	A	A	В	A	В	A	
15	13	15	CRY2_		0		MN1_494		5 1			0.494205		5	A	A	A	A	A	A	A	A
16	14	16	CRY2		1		MN1 59	2863	3 1			0.592867		7	A	A	A	A	A	В	A	A
	15	17	MASC0	7	2		MN1 59					-	9298		Α	В	Α	Α	В	Α	Α	Α
	16	18	MASC0	36								-			В	Α	Α	Α	Α	В	Α	Α
		19	MN1_12	90	3		BKN1			1		-	6158		Α	Α	Α	Α	Α	В	В	Α
		20	MN1_13		4	1	MN1_104	12427	7	1		1.0	4242	В	Α	Α	Α	Α	-	В	Α	Α
		21	AXR1_		5		CRY2_1	021		1	1.187		8784	1	Α	Α	Α	Α	Α	В	Α	В
		22	MASC0	7 1	6		CRY2	429		1		1.1	8843	3	Α	Α	В	-	Α	Α	Α	Α
				1	7		MASC0			1		1 1	8937	4								
							1417 13000			•		1.1	0001	*								

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"description": "Arabidopsis MAGIC data, Gnan et al (2014)",
"crosstype": "magic19",
"sep": ",",
"na.strings": ["-", "NA"],
"comment.char": "#",
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"pmap": "arabmagic_pmap_tair9.csv",
"pheno": "arabmagic_pheno.csv",
"genotypes":
 "A": 1
 "H": 2
 "B": 3
},
"geno_transposed": true,
"founder_geno_transposed": true
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"genotypes":
 "A": 1
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},
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#### Reading data into R

```
library(qt12)
arab <- read_cross2("arab_magic.json")</pre>
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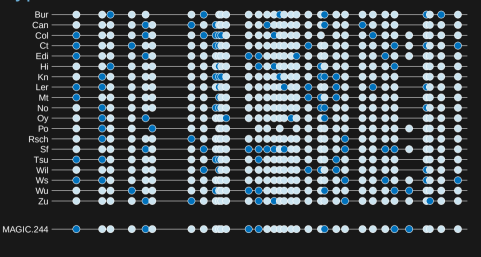
19-way Arabidopsis MAGIC Kover et al. (2009) PLoS Genet Gnan et al. (2014) Genetics github.com/rqt1/qt12data

## Data diagnostics

See Broman et al. (2019) Cleaning genotype data from Diversity Outbred mice. G3 9:1571–1579

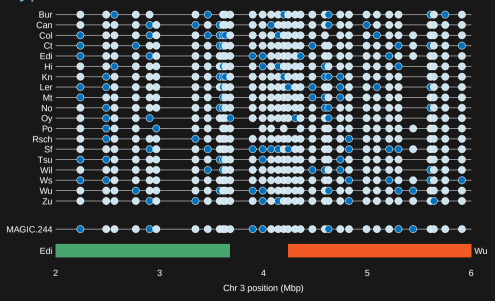
doi: 10.1534/g3.119.400165

#### Genotype reconstruction



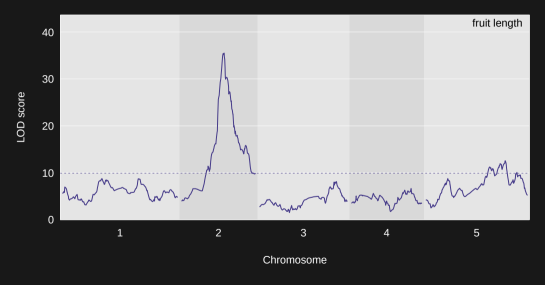
2 3 4 5 6
Chr 3 position (Mbp)

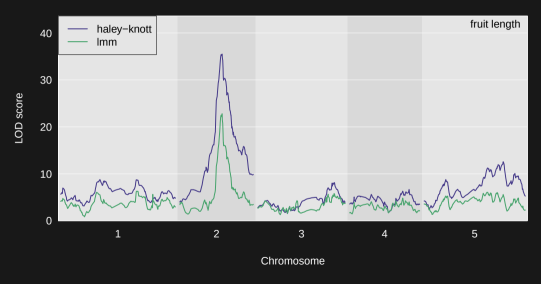
#### Genotype reconstruction

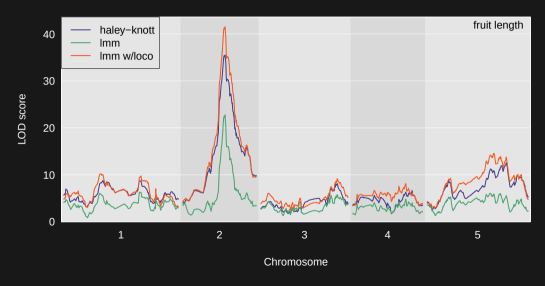


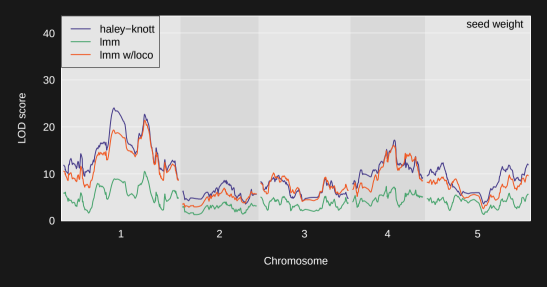
## Genotype reconstruction

```
gmap <- insert_pseudomarkers(arab$gmap, step=0.2, stepwidth="max")
pmap <- interp_map(gmap, arab$gmap, arab$pmap)
pr <- calc_genoprob(arab, gmap, error_prob=0.002, cores=24)</pre>
```

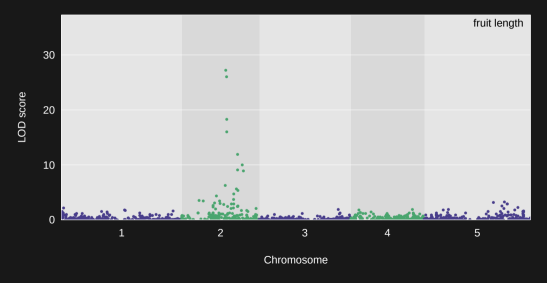


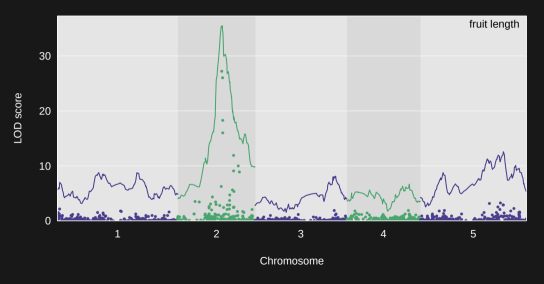


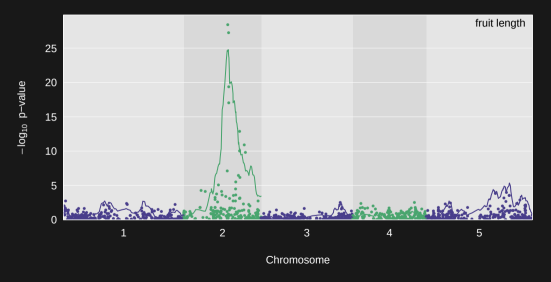


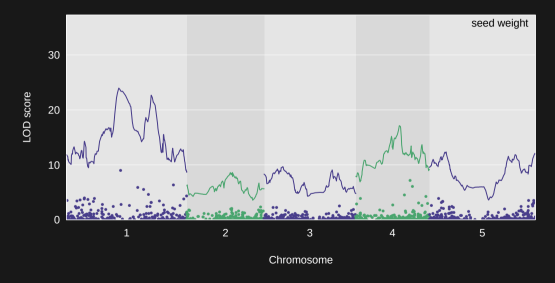


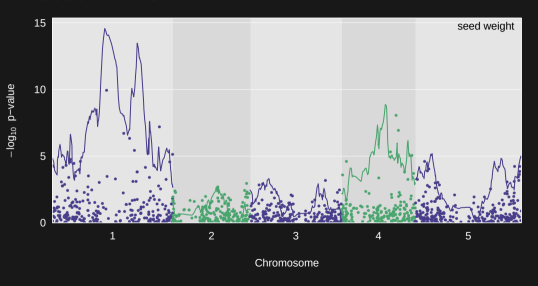
```
out_hk <- scan1(pr, arab$pheno, cores=24)
operm_hk <- scan1perm(pr, arab$pheno, n_perm=1000, cores=24)
k <- calc_kinship(pr, cores=24)
out_lmm <- scan1(pr, arab$pheno, k, cores=24)
k_loco <- calc_kinship(pr, "loco", cores=24)
out_loco <- scan1(pr, arab$pheno, k_loco, cores=24)</pre>
```





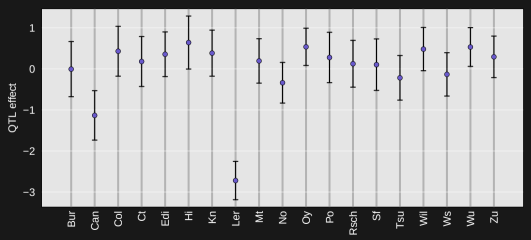




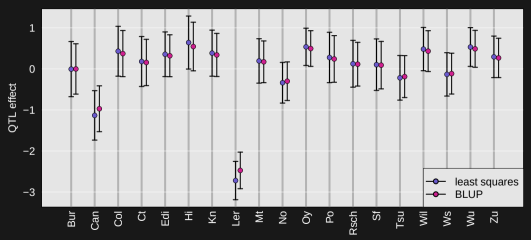


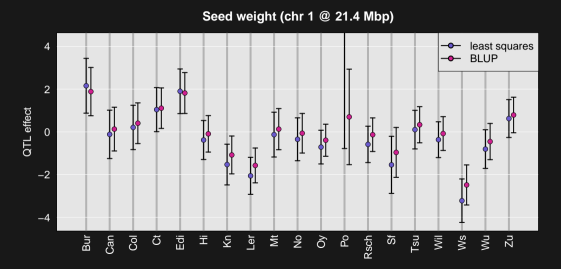
```
snp_pr <- genoprob_to_snpprob(pr, arab)
out_snps <- scan1(snp_pr, arab$fruit, cores=24)</pre>
```











```
fl_peak <- max(out_hk, pmap, lodcolumn="fruit_length")
fl_pr <- pull_genoprobpos(pr, pmap, fl_peak$chr, fl_peak$pos)

fl_fit1 <- fit1(fl_pr, arab$pheno[,"fruit_length"])
fl_blup <- fit1(fl_pr, arab$pheno[,"fruit_length"], blup=TRUE)</pre>
```

#### Goals

- Genotype reconstructions from external software
- ► General models for RIL and AIL
- Sequencing-based genotype data
- Multiple-QTL models
- ► QTL × environment interactions
- ► Interactive data visualization

# Slides: kbroman.org/Talk\_MAGIC2021



kbroman.org

kbroman.org/qt12

github.com/kbroman

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