



MODULE 1 | GENETIC MAPPING

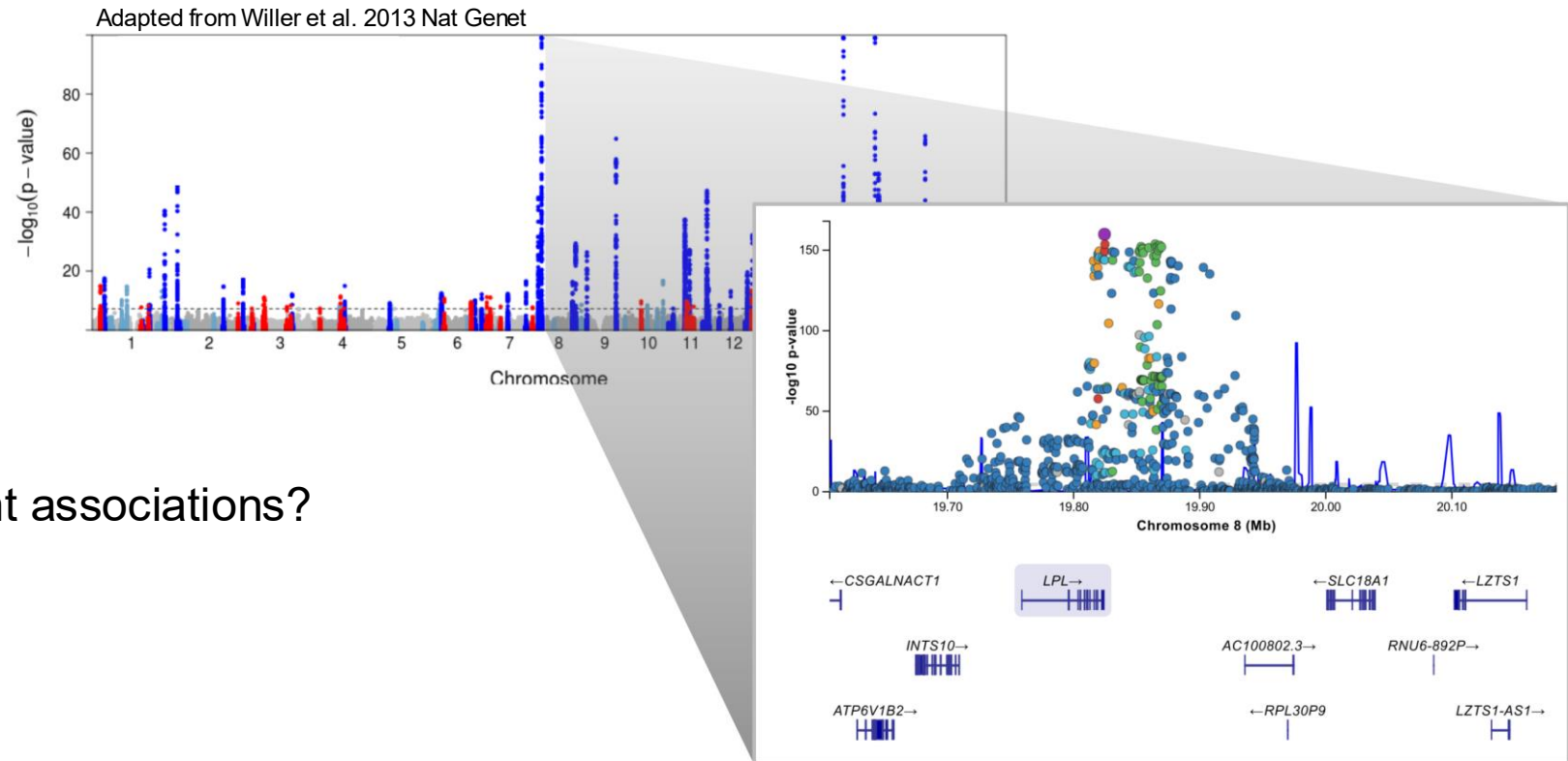
Session 8. Independent GWAS associations

8th July 2025

Slides, practicals & data can be downloaded from the cluster:
`/data/module1/downloadsTuesPM.zip`

Slides and prac guides can be downloaded from the website:
<https://cnsgenomics.com/data/teaching/GNGWS26/module1/>

- GWAS associations are typically clustered along the chromosome
- These clusters typically represent several variants in LD with each other
- How can we identify independent associations?
 - LD-based clumping
 - Conditional analyses
- Why is it important to assess the number of independent associations?
 - Better understanding of the true genetic architecture of the trait
 - Inform functional follow-up studies
 - Out-of-sample prediction

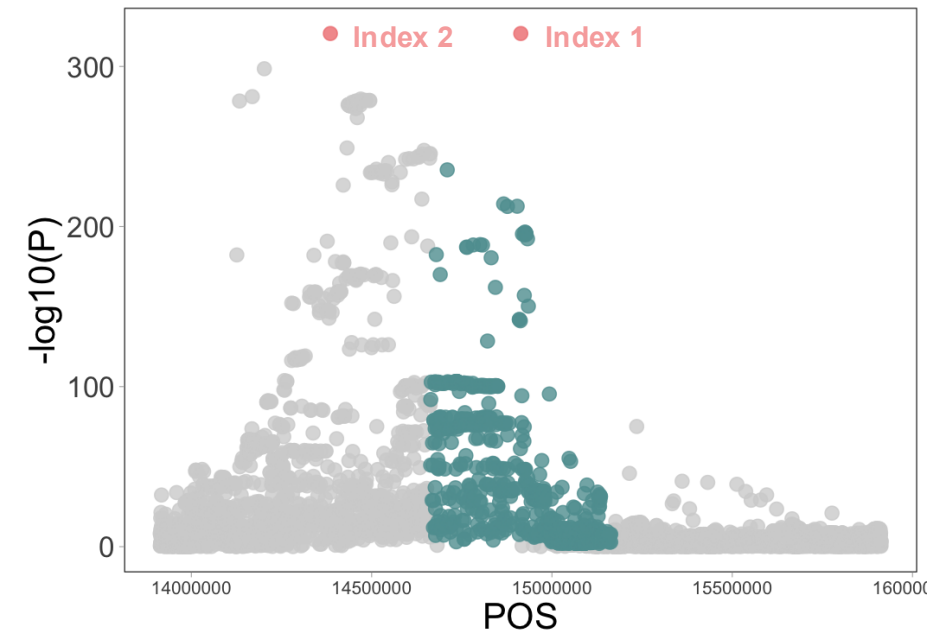


What is LD clumping?

- A method to restrict GWAS summary statistics to a subset of variants that are uncorrelated with each other.
- This method is implemented in PLINK

How does it work?

1. The SNP with lowest P -value is defined as an index SNP
2. Variants are assigned to the same cluster as the index SNP if:
 - in LD with the index SNP, based on a given LD r^2 and associated at a given P -value threshold
 - within a certain distance from the index SNP
3. Start again by assigning next (non-clumped) lowest P -value SNP as an index SNP



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PLINK example:

```
plink --bfile $ld_ref \  
      --clump $sumstats_file \  
      --clump-p1 5e-8 \  
      --clump-p2 0.01 \  
      --clump-r2 0.5 \  
      --clump-kb 250 \  
      --out $output_file
```

Needs individual-level data from the GWAS or a reference dataset to estimate LD

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User can specify:

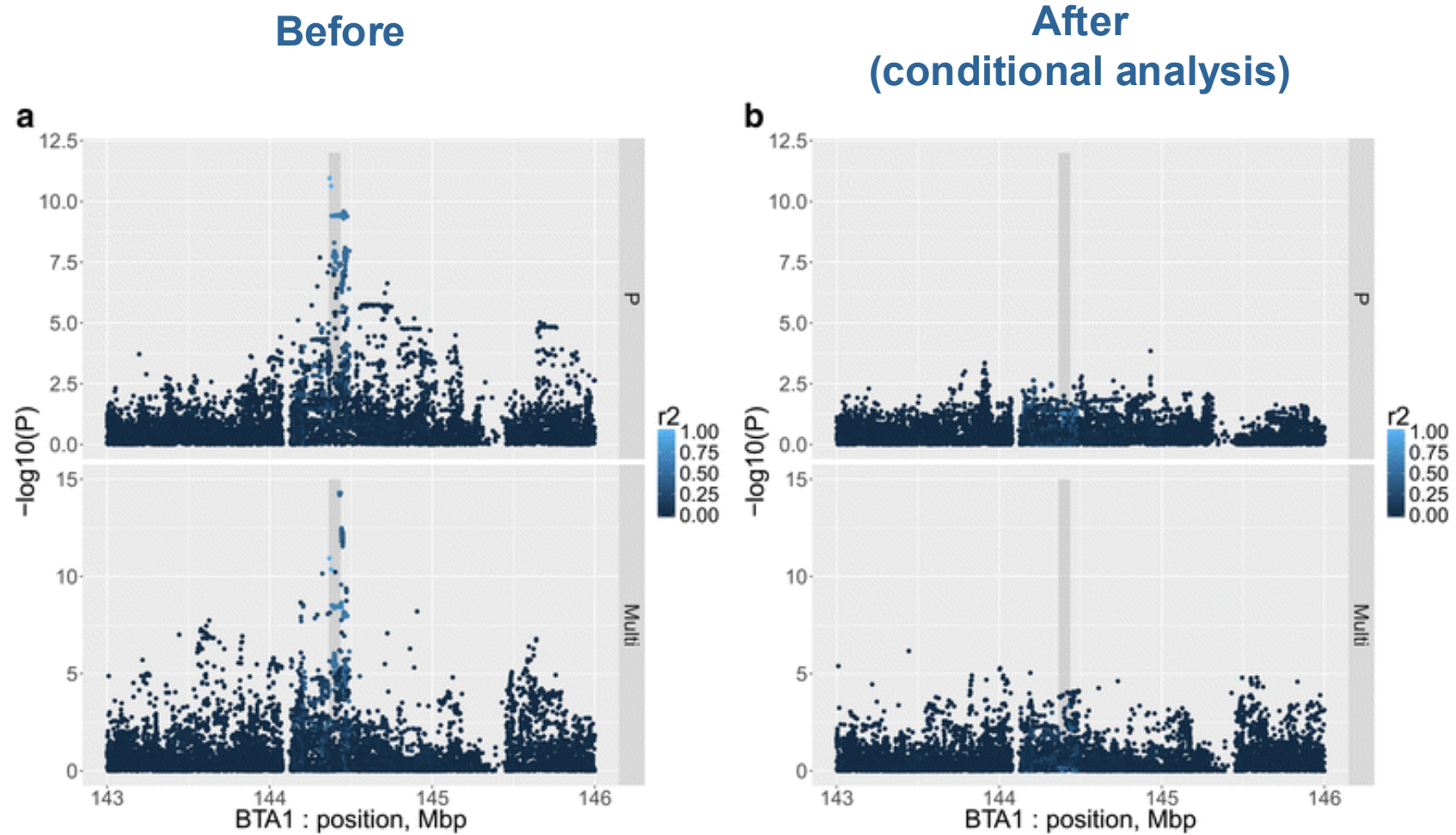
- Maximum P -value of index ($p1$)
- Maximum P -value for clumping ($p2$)
- LD r^2 threshold for clumping
- Distance from index for clumping

Example output file

Each line represents one clump.

CHR	F	SNP	BP	P	TOTAL	NSIG	S05	S01	S001	S0001	SP2
22	1	rs4821083	33056341	2.81e-36	16	0	0	0	0	16	rs762899(1),rs437300i
22	1	rs6007043	45838646	6.75e-32	14	0	0	0	0	14	rs9614466(1),rs214266
22	1	rs7291090	33108150	4.31e-31	9	0	0	0	0	9	rs1475979(1),rs169916
22	1	rs7286215	28397709	3.01e-25	0	0	0	0	0	0	NONE

Clump index SNP (points to SNP column)
Number of SNPs clumped with index SNP (points to TOTAL column)
Number of clumped SNPs with $P < 0.0001$ (points to S0001 column)
Chromosome, base pair and P -value of index SNP (points to CHR, BP, and P columns)
List of SNPs clumped with index SNP (points to SP2 column)



Kemper et al. 2016 BMC Genomics

What do you conclude?

In some cases, we may have variants that are in close proximity and in LD with each other, but that represent independent associations. Here is an example:

Chr	SNP	Position	EA	EAF	beta	P-value	Conditional on rs12794714		Conditional on rs116970203	
							bC	pC	bC	pC
11	rs61883501	13882754	A	0.97	0.002	0.7492	-0.012	0.0373	0.006	0.2683
11	rs116970203	14876718	G	0.97	0.377	0.0E+00	0.415	0.0E+00	NA	NA
11	rs117576073	14912573	G	0.99	0.147	5.9E-62	0.182	1.1E-93	0.159	2.8E-72
11	rs12794714	14913575	G	0.58	0.088	0.0E+00	NA	NA	0.105	0.0E+00
11	rs61891388	66079818	T	0.54	-0.013	4.1E-10	-0.013	4.1E-10	-0.013	4.1E-10
11	rs78168201	70971149	C	0.99	-0.089	5.2E-25	-0.089	5.3E-25	-0.089	5.3E-25
11	rs1660839	71094232	G	0.75	-0.029	6.4E-37	-0.029	6.6E-37	-0.029	6.6E-37
11	rs12803256	71132868	A	0.22	-0.104	0.0E+00	-0.104	0.0E+00	-0.104	0.0E+00

Revez et al. 2020 Nat Commun

- The highlighted SNPs would likely be clumped together because they are:
 - within 36,857 bp of each other and in high LD
- But, conditional analysis shows that the effects are independent
- Ideally – we would conduct a conditional analysis within the sample, but COJO offers a convenient way to do this with summary statistics and an LD reference

Stepwise conditional analysis

An alternative to clumping to identify independent associations is through stepwise conditional analysis. This is typically done with individual-level data, but GCTA implements a method (COJO) that can do these analyses based on summary-level data and an LD reference (Yang et al. 2012 Nat Genet).

How does COJO work?

1. Identify the most significant GWS SNP in the GWAS
2. Calculate the P -values of all other SNPs conditional on the SNP above
3. Identify the next most significant GWS SNP based on the conditional P -values
4. Fit all the selected SNPs jointly in a model and drop the SNP with the largest P -value that is greater than the cutoff P -value.
5. Repeat processes (2), (3) and (4) until no SNPs can be added or removed from the model

GCTA example:

```
gcta --bfile $ld_ref \  
     --cojo-file $sumstats \  
     --cojo-p 5e-8 \  
     --cojo-slct \  
     --out Height_meta
```

User can change specify maximum P -value of SNPs selected to condition on

Example output file

Each line is an independent locus identified with COJO

Info about genetic variant tested				Marginal effects (before COJO) Each variant fitted individually in the model					Joint effects (after COJO) All variants fitted jointly in the same model				
Chr	SNP	bp	refA	freq	b	se	p	n	freq_genos	bJ	bJ_se	pJ	LD_r
22	rs2540653	18973549	G	0.9089	-0.0087	0.0029	0.0029	707907	0.912525	-0.0398	0.003	6.62E-41	0.0034
22	rs1155419	20787850	T	0.5695	-0.0133	0.0017	6.50E-15	700238	0.536779	-0.0265	0.0017	2.35E-53	0.0487
22	rs5754102	21916272	C	0.8226	0.0121	0.0022	4.77E-08	697485	0.824056	0.0317	0.0022	3.36E-45	-0.0342

Effect allele
Estimates are relative to this allele



GWAS summary statistics

Some examples with COJO

SNPs in modest LD

SNP	Chr	bp	Gene	refA	Marginal effects (before COJO)			Joint effects (after COJO)			LD_r
					freq	b	p	freq_gen0	bJ	pJ	
rs17720281	4	145763226	HHIP	T	0.406	0.047	1.70E-30	0.437	0.031	9.20E-13	0.389
rs7689420	4	145787802	HHIP	C	0.837	0.069	1.10E-41	0.836	0.054	2.90E-23	

- marginal SNP effects are overestimated

SNP	Chr	bp	Gene	refA	freq	b	p	freq_gen0	bJ	pJ	LD_r
rs1367226	2	55943044	EFEMP1	A	0.434	-0.005	2.00E-01	0.428	-0.027	5.00E-11	-0.421
rs3791675	2	55964813	EFEMP1	T	0.234	-0.050	1.10E-28	0.249	-0.063	3.00E-37	

- marginal SNP effects are underestimated, a joint analysis is more powerful!

Some examples with COJO

SNP	Chr	bp	Gene	refA	Marginal effects (before COJO)			Joint effects (after COJO)		
					freq	b	p	freq_genot	b _J	p _J
s4932429	15	87164536	ACAN	C	0.51	-0.025	2.70E-10	0.52	-0.022	2.30E-08
rs16942341	15	87189909	ACAN	T	0.031	-0.134	2.20E-24	0.027	-0.116	1.80E-18
rs2280470	15	87196630	ACAN	A	0.334	0.039	1.50E-22	0.33	0.036	6.30E-19

- Multiple significant variants within a locus

Links for further information

- [PLINK clump](#)
- [Yang et al. 2012 Nat Genet](#)
- [GCTA COJO](#)