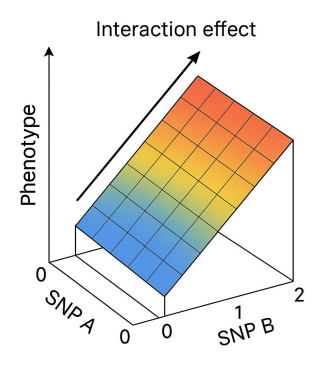
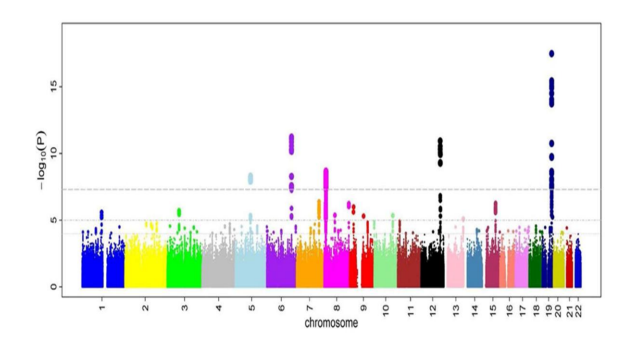
FIGHI: Fisher Information Guided Hyper-Interaction Inference

A scalable and interpretable framework for discovering high-order genomic interactions

Why FIGHI?



- Traditional GWAS misses interaction effects (epistasis)
- Traditional GWAS tests each SNP independently one by one — for its association with a phenotype. That's fine for additive effects, but what if SNP A and SNP B only matter together?



The Challenge: The Combinatorial Explosion

Problem:

P(y|X) may depend on interactions among SNPs.

But enumerating all possible interactions is **combinatorially explosive**:

- With 1M SNPs, even pairwiseinteractions = binom{10^6}{2} → impossible.
- We need a smarter way to decide *which* interactions are worth testing.

So, we reformulate the question as:

"Which combination of SNPs adds the most Fisher Information about the phenotype?"

From Likelihood to Fisher Information

$$\ell(\beta) = \log P(y|X,\beta)$$

Then recall the **score** and **information** definitions:

$$U(eta) = rac{\partial \ell}{\partial eta}, \qquad I(eta) = -\mathbb{E} \left[rac{\partial^2 \ell}{\partial eta^2}
ight].$$

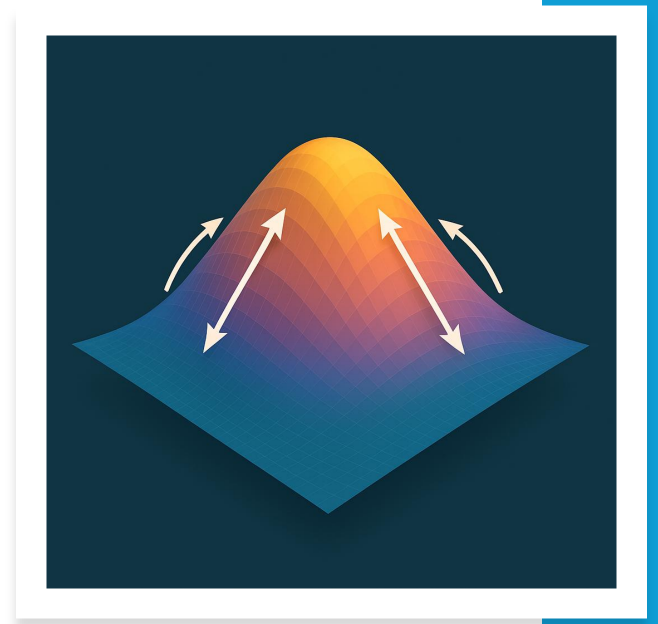
Interpretation:

- $U(\beta)$ tells how the log-likelihood changes with β (slope).
- $I(\beta)$ tells how *stable* that slope is; the curvature, or certainty.
- Large Fisher Information → steeper curvature → more certainty → stronger signal.

So, Fisher Information quantifies how much certainty a parameter contributes.

Intuitive Visualization

Geometric View of Fisher Information



Information Gain for a New Feature

Now suppose we've fit a base model with existing SNPs X, and we're considering adding a new feature:

$$z = x_{j_1} \times x_{j_2} \times \cdots \times x_{j_K},$$

"Potential K-way interaction"

We don't want to refit the whole model for every possible z. Instead, we use the **Score Test Approximation**.

Conceptual Overview

Problem Statement

Given genotype (or multi-omics) data $X \in \mathbb{R}^{N \times P}$ and phenotype y, we seek to identify and rank higher-order combinations of features $S \subset \{1,\dots,P\}$ that contribute synergistically to trait variation.

For each subset S, define an interaction term:

$$\phi_S(x) = \prod_{j \in S} (z_j - ar{z}_j),$$

where z_j is an additive-coded SNP (0,1,2) or a standardized gene/protein score.

We test whether adding ϕ_S to a generalized linear model significantly improves predictive information — quantified by Fisher Information Gain (FIG).

FIGHI Model Structure

FIGHI fits a *progressively expanding* generalized linear model:

$$g(\mathbb{E}[y\mid X]) = lpha + \sum_i eta_i z_i + \sum_{|S|=2} eta_S \phi_S(X) + \sum_{|S|=3} eta_S \phi_S(X) + \cdots$$

but it learns the depth K^* adaptively by tracking information accumulation.

Score Test Derivation

$$U_z = rac{\partial \ell}{\partial eta_z}, \qquad I_{zz} = \mathbb{E} \left[-rac{\partial^2 \ell}{\partial eta_z^2}
ight].$$

From the one-step Newton update:

$$\hat{eta}_{z}^{(1)} = I_{zz}^{-1} U_{z}.$$

Then substitute into the Fisher Information change:

$$\Delta \mathcal{I}(z) = \frac{1}{2} (\hat{eta}_z^{(1)})^2 I_{zz} = \frac{1}{2} \frac{U_z^2}{I_{zz}}.$$

This is the core equation of FIGHI.

Score Test Derivation

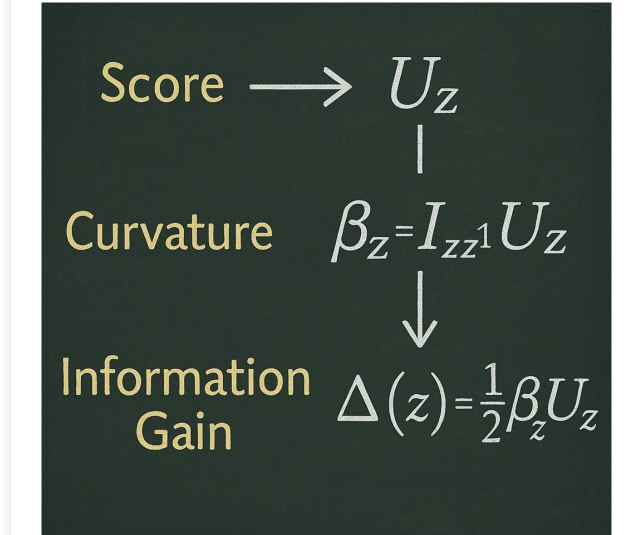
Interpretation

- Uz: how correlated the new interaction is with the residuals
- Izz: how stable (non-collinear) that interaction is
- $\Delta I(z)$ how much extra certainty this interaction adds

So, we can **rank all possible candidate interactions** by $\Delta I(z)$ without refitting the model for each.

Mathematical Consistency

Score-Test Derivation & Computational Simplicity



Fisher Information Gain for Logistic and Linear Models

Logistic case:

We can derive Uz and Izz explicitly.

 $p = \sigma(X\beta), \quad W = \operatorname{diag}(p(1-p)).$

 $U_z = z^ op (y-p), \qquad I_{zz} = z^ op Wz.$

So:

Then:

 $\Delta \mathcal{I}(z) = rac{1}{2} rac{(z^ op (y-p))^2}{z^ op Wz}.$

Fisher Information Gain for Logistic and Linear Models

Linear case:

 $U_z = z^ op (y - Xeta), \qquad I_{zz} = rac{z^ op z}{\sigma^2}.$

We can derive Uz and Izz explicitly.

Hence:

 $\Delta \mathcal{I}(z) = rac{1}{2} rac{(z^ op (y - Xeta))^2}{z^ op z}.$

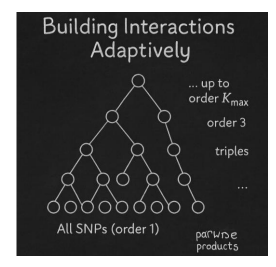
Both cases require only **vector operations** — no full refitting.

Building Interactions Adaptively

But we don't explore everything. We keep only the interactions that **add significant Fisher Information**.

FIGHI **adapts its order depth** — if information saturates early, it stops at 2-or 3-way.

That's why even if we set max_order=4, you may only see 2-way interactions — because that's where Fisher Information stops growing.



Define:

$$r_K = rac{\sum_{k=1}^K \Delta \mathcal{I}_k}{\sum_{k=1}^{K_{max}^{theor}} \Delta \mathcal{I}_k}.$$

When rK > threshold (say 0.95), stop.

Computational Implementation

Memory-efficient pipeline.

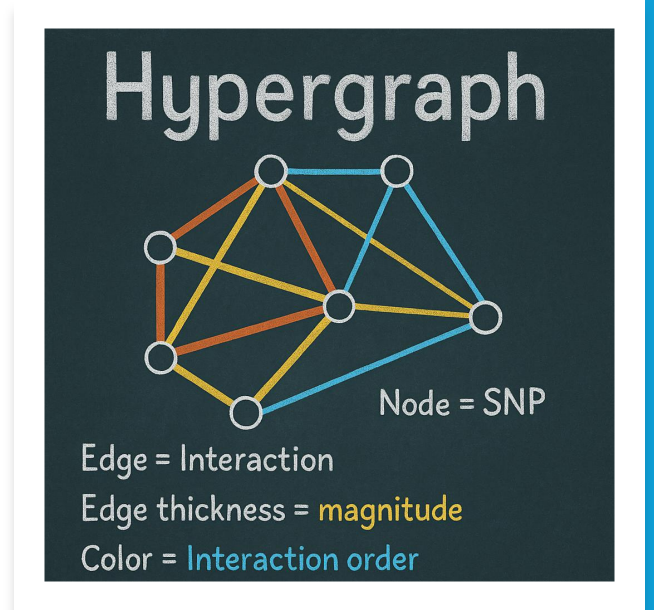
- Reads huge genotype CSVs in chunks (read_csv(chunksize=...))
- Uses **prescreening**: keep only top M SNPs by correlation with phenotype
- Streams data blockwise, computes ΔI(z) incrementally

Hypergraph Representation

Say:

- Each SNP = node
- Each discovered interaction = hyperedge
- Edge weight = $\Delta I(e)$

So, the **hypergraph encodes multi-level cooperation** among SNPs.



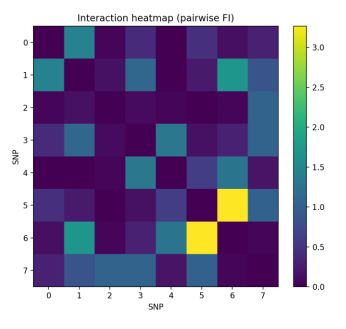
Simulation

1	case	rs101	rs102	rs103	rs104	rs105	rs106	rs107	rs108
2	1	1	1	1	1	1	0	1	0
3	1	1	1	1	1	1	0	0	0
4	0	2	1	0	0	0	0	0	0
5	1	1	1	0	0	1	0	0	0
6	1	0	1	0	1	1	1	1	0
7	1	2	1	1	1	2	0	0	1
8	1	1	1	0	0	1	0	1	0
9	1	1	0	0	0	1	0	1	1
10	0	0	0	0	1	1	0	0	0
11	1	1	1	1	0	1	0	1	1
12	0	1	0	0	1	1	0	1	1
13	1	2	0	2	0	2	0	0	1
14	1	1	1	1	0	1	0	0	0
15	1	2	0	0	1	0	0	0	1
16	0	1	0	1	0	0	0	0	0
17	0	0	0	0	0	1	0	0	0
18	1	1	0	2	1	1	0	1	0
19	0	0	1	0	1	0	0	0	0

Simulation Example

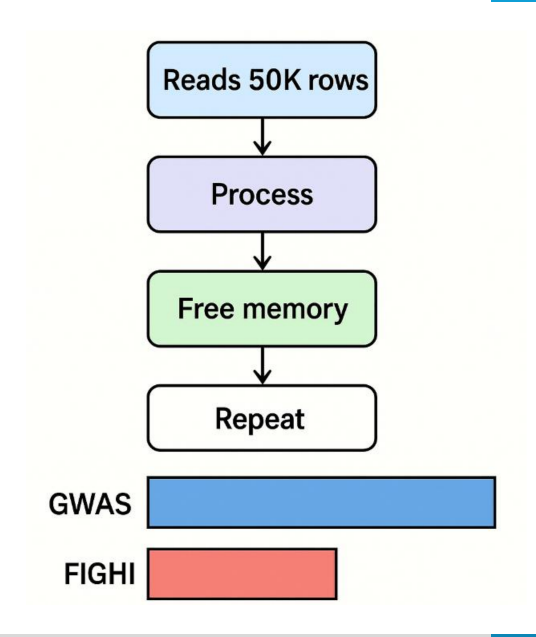
1	hyperedge	order	fi_gain	pval	beta_hat	info
2	rs106 rs107	2	3.265700508971495	nan	0.7577984469105081	11.373625598196089
3	rs102 rs107	2	1.7152008230200526	nan	0.41324759714950715	20.087426376755193
4	rs101 rs102	2	1.4421667058999605	nan	-0.4855839405979781	12.232545236598138
5	rs104 rs105	2	1.290800063735504	nan	0.388608570407242	17.094807912396966
6	rs105 rs107	2	1.2510852582980987	nan	-0.4224610390272857	14.019856768862953
7	rs102 rs104	2	1.071877164735371	nan	-0.31743233505528967	21.275152747506475
8	rs103 rs108	2	1.042533010722737	nan	-0.38027087946879984	14.418951226315185
9	rs104 rs108	2	1.0385913559735784	nan	-0.41622726641193086	11.989847127964518
10	rs106 rs108	2	1.0174236222765263	nan	-0.4854604808314015	8.63423866243303
11	rs102 rs108	2	0.8521801770980068	nan	-0.3679639819301112	12.587828098196317
12	rs105 rs106	2	0.5872604638410686	nan	-0.252451038265352	18.429198045406924
13	rs101 rs106	2	0.4489338152192574	nan	-0.30473903308178546	9.66843416584441
14	rs101 rs104	2	0.4070612959686377	nan	-0.2747933052539285	10.781458383678162
15	rs101 rs108	2	0.30287425528108963	nan	-0.186134667263455	17.48388013488581
16	rs104 rs107	2	0.2962962928135718	nan	-0.20740742281203575	13.775507995883775
17	rs102 rs106	2	0.2411277582608408	nan	-0.16978446685606224	16.72943576626595
18	rs105 rs108	2	0.16676502208568447	nan	-0.14015875807616904	16.9783108131993
19	rs104 rs106	2	0.14613526489890064	nan	0.16030351654085637	11.373625598196089

1	SNP	FI_total	Fl_main	FI_interact	MAF	Rank	Gene	Pathway
2	rs107	3.3766863527319053	0.0	3.3766863527319053	0.25	1		
3	rs106	2.8580899735984246	0.0	2.8580899735984246	0.06666667014360428	2		
4	rs102	2.736520403358848	0.0	2.736520403358848	0.32499998807907104	3		
5	rs108	2.2323323149110132	0.0	2.2323323149110132	0.14166666567325592	4		
6	rs104	2.1706485955217376	0.0	2.1706485955217376	0.25	5		
7	rs105	1.6761473622990806	0.0	1.6761473622990806	0.36666667461395264	6		
8	rs101	1.3899051013540429	0.0	1.3899051013540429	0.4583333432674408	7		
9	rs103	0.7108106009866798	0.0	0.7108106009866798	0.3083333373069763	8		



Computational Efficiency

Memory-efficient pipeline.



Biological and Theoretical Takeaways

- **Biological** Captures epistasis efficiently, beyond GWAS main effects.
- Mathematical Based on score test and Fisher Information curvature.
- Computational Scales to large datasets via streaming and pruning.

$$\Delta \mathcal{I}(z) = rac{1}{2} rac{U_z^2}{I_{zz}}$$

THANK YOU!

Full FIGHI documentation and code:

https://github.com/1234-Ariel-code/fighi