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Wiki page: https://simons-institute.github.io/pseudorandomness/groups/modelstructure.html

1 Big picture

I'll talk about things which have different names in different fields. You probably know them, but don't know they are the same or related ideas that come up in different fields.

	Boosting	Hard-core	Dense model	Weak regular-	?
		lemma	theorem	ity	
Area	Area ML		Additive	Graph theory	
			combinatoric-		
		ization	s/CC		
Credit	Shapiro, Fre-	I, Hol	GT, Barak-	Sz,	
	undS		SWigderson	FriezeKannan	
Get	Circuit com-	"	Proof that set	"	
	puting $f 1 - \delta$		isn't δ -dense		
	of the time				
Unless	Weak nearner	Hard-core	$\Omega(\delta)$ -dense	A model	
	fails on distri-	distribution	"model"	succinctly	
	bution of den-		indistinguish-	describing set	
	sity $\Omega(\delta)$		able from		
			set		
Algorithm	Weak learner	"	Distinguisher	"	
needed to be					
const.					

Arrows:

- 1. KS
- 2. I
- 3. TTV, RTTVadhan

Klivans Servedio.

If set looks like might be dense, either succinct reason (proof) not δ -dense, or dense model (reason why it looks big to certain class of sets)

Hard-core lemma implies dense model theorem.

FK simpler argument with better quantitative bounds, weaker conclusion. Generalized to sparse graphs without giveaway cut (cut prove not dense). Either you get a proof that the cut isn't dense, or get a model that succinctly describes all cuts.

Take a bunch of theorems we know are true and showing implications between them. Hope a lot to gain by looking at connections between them:

1. Versatility. Retrofit. Ex. there are many boosting algorithms. When you follow this progression, you get different quantitative and qualitative versions of dense model theorem and regularity.

Boosting is a particular proof for the min-max theorems.

2. Algorithmic and constructive:

There are nonconstructive versions using the min-max theorem for boosting, hard-core lemma, dense model theorem.

I care about algorithmic versions.

In ML we care about getting a function that computes a function much of the time.

On the other side, we're really after the distribution where the weak learner fails, the model that succinctly describes the set. We pay attention to do the reductions in an algorithmic, not just an existential way.

- 3. Dense model theorem \rightarrow learning. Take boosting technique and use it in an unsupervised way.
- 4. Generality: some things seem to be specific to a setting (density of graphs). Actually weak regularity doesn't have anything to do with graphs being dense. We can relativize it to subgraphs of any graph. You can look at subgraphs of expanders, bipartite graphs, etc. and plug it in the same machineary. If you want to look at spectral norms rather than cuts, you can plug into the same machinery.

Almost all arrows are trivial! Simple lines of high-school algebra.

Weak regularity does imply dense model theorem and hard-core lemma.

1.1 Boosting

First we discuss the PAC learning model.

Let U be the universe—big set of possible inputs. There is a boolean function f and we get points (x, f(x)) where $x \sim U$. The algorithm can request any number of such points. Our goal is to find a hypothesis h such that

$$\mathbb{P}_{x \sim U}[h(x) = f(x)] \ge 1 - \delta.$$

How do we get a hypothesis h? We assume we have a weak learner, which receives the same kind of input, but perhaps from some other distribution μ , and gets something correlated with the right answer.

$$\mathbb{P}_{x \sim \mu}[h_i(x) = f(x)] \ge \frac{1}{2} + \varepsilon.$$

(It has to work on any distribution μ satisfying some assumptions.)

How do we use the weak learner? Give a routine that subsamples the distribution U to pass to μ . We let $\mu: U \to [0,1]$. Keep something in U and keeping it with probability in [0, 1], else throw it back (rejection sampling). This induces a probability distribution

$$D_{\mu}(x) = \frac{\mu(x)}{\sum_{x' \in U} \mu(x')}.$$

Let $|\mu| = \mathbb{E}_{x \in U} \mu(x)$. Use this as a notion of density. Condition:

1. We will only run the weak learner on distributions of about the same density, $|\mu| >$ $\Omega(\delta)$.

We don't want to run it on something too puny, because the time to simulate the distribution is inversely proportional to the density.

2. Each $h_i \in T$. Ex. T is linear separators.

$$F_kT = \{f(h_1, \dots, h_k) : f \in F, h_1, \dots, h_k \in J\}.$$

It turns out that $\mu \in F_KT$: it's basically describable in the same class of functions. $h \in F_K T$.

Pick one large sample, run on it as much as you'd like. Then the issues of what is μ disappears.

If you just care about ML we don't care about this details, but to go further, we do.

The first boosting algorithm I'll give is totally ridiculous from the ML point of view. For people who work on weak regularity on graphs this is the natural version and leads to the standard versions of results.

Suppose you just look at things information-theoretically. If I learn these hypotheses are relevant, how should I use them to predict the function? What's the best function that composes k of these things and gives a predictor for f?

 h_1 is either T or F, etc. This divides into 2^n buckets. Once you are in a bucket what should you do? When given the next x, all you know is which bucket it's in.

To optimize, you predict the majority. Let p_B be the probability of the majority; then $1 - p_B$ is the minority. Let $H_i(x)$ be the majority answer in B. B is distributed according to B(x). It takes 2^i time to give a good estimate for the majority answer.

Let $\delta_i = \mathbb{E}_{B \in B[x], x \in U}[1-p_B]$. If the majority $\to 1$ in every bucket, then our error probability is small. While it's big, we still need to learn.

On the same biased distribution, get the same h_1 . We want to make sure not boolean combination of previous h_i 's is a good predictor for the next distribution. If $f(x) = \mathsf{Maj}_B$, then $\mu(x) = \frac{1-p_B}{p_B}$ for $x \in B$. What is $|\mu|$?

$$\underset{x}{\mathbb{E}} |\mu| = \underset{x}{\mathbb{E}} \mu(x) = \underset{B}{\mathbb{E}} [\underset{x}{\mathbb{E}} [\mu(x)|x \in B]] \tag{1}$$

$$= \mathbb{E}[1 - p_B + 1 - p_B] = 2 \mathbb{E}[1 - p_B] = 2\delta_i. \tag{2}$$

If measure small, then small failure probability. If haven't succeeded, then μ is large enough so we can continue learning on μ .

Given previous functions, define candidate function. Either it has good success, or distribution has good density, and we continue. I need to show this process terminates within a reasonable amount of time.

We'll look at a potential function that measures how close we are to success. Our goal is to drive all p_B 's to 1.

Our potential function is

Flip a coin with expected probability.

$$\varphi_i = \mathop{\mathbb{E}}_{p_0} p_1 (1 - p_1) = \mathop{\mathbb{E}}_{p_0} p_1 - p_1^2 \tag{3}$$

$$= \mathop{\mathbb{E}}_{B} p_{B,1} (1 - p_{B,1}) \tag{4}$$

Land in B_1 with probabilty q, B_0 with probabilty 1-q. f=1, $p_{B,1}+\alpha_+$, $p_{B,1}-\alpha_-$. $q\alpha_t=(1-q)\alpha$.

$$q(p_{B_1} + \alpha_+)^2 + (1 - q)(p_{N,1} - \alpha_-)^2 = (q + 1 - q)p_{B,1}^2 + (\alpha_t q - (1 - q)\alpha - 1)p_{B,1} + q\alpha_t^2 + (1 - q)\alpha_{-i}^2.$$

(5)

$$\mathbb{E}[\mu(x)(-1)^{f(x)\oplus h_{i+1}(x)}] \ge \varepsilon |\mu| \tag{6}$$

$$= \underset{B \in B(x)}{\mathbb{E}} \underset{x \in B}{\mathbb{E}} \mu(x) (-1)^{f(x) \oplus h_{i+1}(x)} \tag{7}$$

$$q\left(\frac{1-p_{B}}{p_{B}}(p_{B}+\alpha_{t})-(1-p_{B}+\alpha_{-})\right)$$
 (8)

$$(1-q)\left(\frac{1-p_B}{p_B}(1-p_B+\alpha_-)-(p_B+...)\right)$$
(9)

Prob h on B is q. For f = 1, $q + \alpha_+$. For f = -1, $q - \alpha_-$. Then

$$p\alpha_+ = (1 - p)\alpha_-.$$

Then

$$\mathbb{E}_{\mu}\mu(x)(-1)^{f(x)\oplus h(x)} = \varepsilon_B = p\frac{1-p}{p}(q+\alpha_+ - (1-q-\alpha_+))...$$
 (10)

$$(1 - q - \alpha_{-}) - (1 - \alpha_{-}) \tag{11}$$

$$(1-q)(2\alpha_{+} + 2\alpha_{-}). (12)$$

On the other hand the expectation

$$p_{B,1} = q \left(\frac{p(q + \alpha_+)}{q}\right)^2 - (1 - q) \left(\frac{p(1 - q - \alpha)}{1 - q}\right)^2$$
(13)

$$=p^2\left(q+\frac{2\alpha_+}{q}+\frac{\alpha_+^2}{q}\right)()\tag{14}$$

Try to make an energy increment argument.

It matters that the weak learner only looks at distributions with density $\geq 2\delta$. From boosting to hard-core:

Lemma 1.1. Let f be any function on U and T any class of tests. Either there exists $H \in F_K T$, H(x) = f(x) with probability $1 - \delta$, or there exists μ , $|\mu| \ge 2\delta$, $\forall h \in T$,

$$\mathbb{P}_{x \in \mu}[h(x) = f(x)] \le \frac{1}{2} + \varepsilon,$$

$$k = O\left(\frac{1}{\varepsilon^2 \delta^2}\right)$$
.

Proof. Let weak learner be exhaustive search. get sequence of distribution, all of $\geq 2\delta$. See if any of them have a good bias. If we always get something with bias $\geq \frac{1}{2}$, continue in the strong learner, until we get H(x) = f(x) with probability $1 - \delta$. If at some point our exhaustive search algorithm gets stuck, we have a distribution that's hard core.

Region where very hard, indistinguishable from random.

Why have to keep track: we want size of hard-core to be what it should be. 2δ is what it should be.

A lot of times we want to compute one or the other. Then we need to use the boosting algorithm. By using the boosting algorithm we get something stronger. We get that the hardcore distribution doesn't have unlimited complexity. It's describable by the function we want to compute, $\mu \in F_{k+1}(J \cup \{f\})$.

Let $S \subseteq U$. We have some tests T that don't reveal that it's small. If big, then δ fraction of whole thing.

Definition 1.2: Let $S \subseteq U$. Let \mathcal{T} be a class of tests. For $T \in \mathcal{T}$, $\mathbb{E}_{x \in S} T(x)$. S is (ε, δ) -pseudo-dense for \mathcal{T} if for all $T \in J$, $T(U) \geq \delta T(S) - \varepsilon$. This is written as $S \sim_{\varepsilon, \delta} U$.

One way to be pseudo-dense it to be dense. Another, one step removed, is that there's R that's indistinguishable from the whole distribution, and it's dense in R. and T(U) and T(R) are within δ fraction of each other.

Theorem 1.3 (Dense model theorem). There exists μ , $|\mu| \geq \delta$ and $S \sim_{\varepsilon,,\delta} D_{\mu}$. If S is (ε, δ) -pseudodense vs $F_k \mathcal{T}$, $k = O\left(\frac{1}{\varepsilon^2 \delta^2}\right)$ then there exists μ , $\mu \in F_k \mathcal{T}$, $|\mu| \geq \delta - \varepsilon$, $D_{\mu} \sim_{\mathcal{T}} S$ to within $O(\varepsilon/\delta)$.

By going through boosting and hard-core, we get characterization of μ , describable using original tests.

Hard function should be membership in S. This happens almost never, so we have to blow up membership to make it comparable to what it should be.

Proof. Let U' be: δ' fraction is $(0, x), x \in S$, while $1 - \delta'$ is $(1, x), x \in U$. Let $\delta' = \frac{\delta}{1 + \delta}$. Let f(b, x) = b, T((b, x)) = f(b, x) with probability $1 - \delta'$.

The tests are just applied to the x part.

$$\delta'T(S) + (1 - \delta')(1 - T(U)) = 1 - \delta' + \delta'(T(S)) - (1 - \delta')T(U)$$
(15)

$$=1-\delta'+\frac{1}{1+\delta}(\delta T(S)-T(U))\leq 1-\delta'+\varepsilon. \tag{16}$$

This distribution has a hardcore subdistribution on which is as hard as a random coin flip. With δ' probability, sample from S, f=1. with $1-\delta'$ probability, sample from U, f=0. The subset has size $2\delta'$, in order to be hard, half comes from S (all of it), half from U. f hard to predict means the 2 sides indistinguishable.

The model depends on k and f. Magically when we apply it we only want the part of the model where f = 0. It's definable with k things from \mathcal{T} . No membership oracle required. \square

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I'll start with motivation: one thing we could hope to get from this.

1. Let X be a set in \mathbb{R}^n

Let S be some distribution on points in X.

Let J be a set of classifiers, ex. a set of half-planes.

Let F_KJ be boolean functions on K functions in J; here, partitions into polygonal regions by k half-planes.

We want to pre-process the distribution to be able to answer queries in F_KJ .

- 2. Here, violation of pseudo-density means: a polygonal region with many more points from S than its volume, a "hot spot".
- 3. A model is a partition into polygonal region, with probability distribution on regions. Estimate half-space probabilities by as if uniform within regions.

A simple model is defined by at most k lines. Uniformly sample within region.

$$Area (region) < \delta \mathbb{P}_S (region) - \varepsilon.$$

"Given any half-space, how many points are on one side of it?"

4. Algorithmic requirement: given points from subsample of S, points from subsample of U, find a half-space that approximately maximizes the difference in red probabilities and blue probabilities.

When using boosting, the equivalent is a distinguishing algorithm. Find a separating halfspace that approximately maximizes the probability distribution under the two distributions.

"Using boosting in unsupervised context": not told what we want to learn about distribution. Just have model to address queries later on. You need tests in mind.

Setting	Boosting	Hard-
	WL: $ \mu_i \geq 2\delta, \mu_i = g(h_1, \dots, h_K, f), h_{i+1} \in \mathcal{T}, K$ iterations	Hardcore measure: μ_K
	SL: $H = G(h_1, \dots, h_K), \mathbb{P}[H = f] \ge 1 - \delta$	Violation of hardness: $H =$
Assumption	WL never fails	Violatio
Conclusion	SL works	Hard-core meas
Algorithmic	Weak learner requirement	Approximately

Weak regularity: Same as above Assumptions: Actually dense Conclusion: model exists

Notes:

1. Boosting: δ is the same. Many boosting algorithms meet this criterion.

"Either weak learner fails or strong learner works." In boosting we think of weak learner as never failing.

2. For hard-core measure lemma: we want to come up with the measure. Either we can find hard-core measure: no compute more than $\frac{1}{2} + \delta$ of time, or find a function computing $1 - \delta$ of time.

The only change is psychological: we assume violations never happen. The logical format is the same.

3. Every boosting gives hard-core measure lemma. It gives a hard-core measure lemma with the same parameters. The way we glue the functions is exactly the same.

We have different hard-core measure lemmas with different gluing. Sometime you care about computational complexity of G but not of g, etc. Or it could be the other way around.

- 4. Convert to dense model theorem/transference principle (Tao and Ziegler). Either we have a distribution we're trying to model, distribution has pseudo-density property: there won't be a violation that's definable from K different properties from hypothesis class, where violation means that it's much smaller on U than on S. We get model of density 2δ . Get model definable in terms of K hypotheses from class. Assume violation of pseudo-density does not happen, then we get a model.
- 5. Weak regularity: is just DMT except the distribution actually is dense. It's not so interesting that it has a dense model. But what this says is that the dense model you get is simple, definable in terms of a small number of basic hypotheses.

applying to actually dense, so you don't worry about violation of pseudo-density.

Sometimes we care about simplicity in model, sometimes in G.

6. Note the k is the same. Reductions preserves k, and the functions h_i, G . When I talk about an implication, not jsut known boosting gives known hard-core lemma, regularity lemma, I mean whatever boosting algorithm you give me, it gives me a

hard-core lemma, regularity lemma with the same parameters. We can immediately jump to whatever we're interested in. Pick the one that gives the best results.

Boosting with oblivious decision trees.

Let

- f be function we want to learn, $f: \chi \to \{0, 1\}$.
- \bullet \mathcal{T} class of Boolean functions we use as basic hypotheses.
- $F_k \mathcal{T}$ compositions of $\leq k$ functions in \mathcal{T} .

Measure $\mu: X \to [0,1]$, $|\mu| = \mathbb{E}_{x \in X} \mu(x)$, $D_{\mu}(x) = \frac{\mu(x)}{|\mu||X|}$ induced distribution: pick x, keep with probability $\mu(x)$, else repeat.

- Weak learner: If $\mu \in F_k \mathcal{T}$ and $|\mu| \geq 2\delta$, who returns $h \in \mathcal{J}$ so that $\mathbb{P}_{x \sim D_{\mu}}[h(x) = f(x)] \geq \frac{1}{2} + \varepsilon$.
- Strong learner: whp, finds $H \in F_k \mathcal{T}$, $\mathbb{P}_{x \in X}(H(x) = f(x)) \ge 1 \delta$.

Let μ_0 be constant 1. While $|\mu_i| \ge 2\delta$ do

- $h_{i+1} \leftarrow WL(\mu_i)$
- Partition X according to values of h_1, \ldots, h_i into "blocks" B, Maj_B most likely value in δ .
- $H_{i+1} = \text{output Maj}_{B(x)}$,

$$\mu_{i+1} = \begin{cases} \frac{1 - p_{\mathsf{Maj}}}{p_{\mathsf{Maj}}}, & \text{if } f(x) = \mathsf{Maj} \\ 1, & \text{otherwise} \end{cases}$$

cost exponential in k in most circumstances. under what circumstances not exp in k? If VC dimension small.

number of boolean comb. Grows at most order k^d .

Induce new distribution, make f unbiased within each block.

Total probability: on $1 - p_{Maj}$, f = 1, make p_{Maj} . f = 0, $1 - p_{Maj}$.

(Last time: if $|\mu_i| \le 2\delta$, $\mathbb{P}_{x \in X}[H_i = f] \ge 1 - \delta$.)

We need to show this method terminates.

Consider the potential function

$$\varphi = \mathbb{E}_{x \in UX, B \leftarrow B(x)}[(\mathbb{P}[f=1|B])^2]$$

Closer to 1, closer it is to boolean. This is maximized if every bucket/block is either constantly 1 or 0.

We need to show every iteration increases this potential function.

WTS: In every iteration φ increases by $\Omega((\varepsilon\delta)^2)$. Fix B.

$$p = \mathbb{P}[f = 1|B] \tag{17}$$

$$q = \mathbb{P}[h_{i+1} = 1|B] \tag{18}$$

$$q + \alpha_{+} = \mathbb{P}[h_{i+1} = 1|B, f = 1] \tag{19}$$

$$q - \alpha_{-} = \mathbb{P}[h_{i+1} = 1|B, f = 0] \tag{20}$$

$$\alpha_{+}p = \alpha_{-}(1-p)$$
 by conservation (21)

$$B \leftarrow \alpha_{-}(1-p) \tag{22}$$

$$p_0 = \mathbb{P}[f = 1|B_0] = \frac{\mathbb{P}[f = 1 \land h = 0]}{\mathbb{P}[h = 0]} = \frac{p(1 - q - \alpha_+)}{1 - q}$$
(23)

$$p_1 = \mathbb{P}[f = 1|B_1] = \frac{\mathbb{P}[f = 1 \land h = 1]}{\mathbb{P}[h = 1]} = \frac{p(q + \alpha_+)}{q}$$
 (24)

(Value of potential) =
$$qp_1^2 + (1-q)p_0^2 = p^2 \left(\frac{(q+\alpha_+)^2}{q} + \frac{(1-q-\alpha_+)^2}{1-q} \right)$$
 (25)

$$= p^{2} \left(\left(q + 2\alpha_{+} + \frac{\alpha_{+}^{2}}{q} \right) + \left(1 - q - 2\alpha_{+} + \frac{\alpha_{+}^{2}}{1 - q} \right) \right)$$
 (26)

$$= p^2 \left(1 + \frac{\alpha_+^2}{a} + \frac{\alpha_+^2}{1 - a} \right) \tag{27}$$

$$\geq p^2 + 4p^2\alpha_+^2 \geq \alpha_+^2 \tag{28}$$

if Maj = 1.

Assume $Maj_B = 1$.

$$\underset{x \in B}{\mathbb{E}} [\mu(x)((-1)^{(h(x) \neq f(x))})] = (f = 1)p\left(\frac{1-p}{p}\right)[(q+\alpha_+) - (1-q-\alpha_+)]$$
(29)

$$+ (f = 0)(1 - p)1[1 - (1 - \alpha_{-}) - (q - \alpha_{-})]$$
(30)

$$= (1-p)[2\eta - 1 + 1 - 2\eta + 2\alpha_{+} + 2\alpha_{-}] + 2\alpha_{+}(1-p) + 2\alpha_{+}p = 2\alpha_{+}$$
(31)

$$\mathbb{E}_{x \in B} \dots = \mathbb{E}_{x} \mathbb{E}_{B = B(x)} \mu(x) (-1)^{(f(x) \neq h_{i+1}(x))}$$
(32)

$$= \mathop{\mathbb{E}}_{B} 2\alpha_{+} \ge \varepsilon |\mu| = 2\delta \varepsilon \tag{33}$$

$$\Delta \varphi = \mathop{\mathbb{E}}_{R} \alpha_t^2 \ge (\delta \varepsilon)^2. \tag{34}$$

Every step don't terminate, potential function increases by $(\delta \varepsilon)^2$. It's a number in [0,1], so the number of iterations is at most $k \leq O\left(\frac{1}{(\delta \varepsilon)^2}\right)$. At most polynomial in parameters.

All you get from this is a decision tree. Brute force. Complexity is exponential in k. This is a bug, not a feature.

Can always convert sign into decision tree. On hard-core, DMT give same thing. Incomplexity terms, we don't get good hard-core, because circuit size 2^k circuit size. Better boosting algorithm makes G have smaller complexity. If stopping point is HCL, not boosting

you want. In dense model theorem, fine because all you care about is size of k not complexity of G.

This potential function matches this boosting algorithm. Other boosting algorithms you can analyze with other potential functions.

Why I want to do this way with this potential function:

- Like potential function used most in graph theory. Property: you can't make negative progress, you always go forwards.
- Translated get usual Sz, except don't get equal size partitions.

We have different batches/blocks that we're partitioning to. Suppose we get stuck: no function that correlates globally, but there are many blocks where we can find functions that correlate with the function inside that block. If ε fraction of blocks find functions that correlate, query them all. That's going to be our next round.

 2^k functions correlating. Now 2^{2^k} buckets, but increased potential function by poly in terms of ε , δ . This is a familiar argument, we can only go $\frac{1}{\varepsilon}$ iterations before we terminate.

We've increased everything a tower depending on ε . ε had better be a big constant. We have a hard-core defined in terms of k things. Hard-core looks like: partition. $1 - \delta$ fraction where have answer correct. Boolean combinations of k things you've learned. Except for ε fraction, each part: no better method than predict constant function.

Terminate: on most blocks we can't beat majority answer.

Bounds: you need a tower. No avoidance of tower in some contexts. Graphs are not the same context.

strong hardcore, or strong dense model. I don't know the formulation let alone the proof. I have the proof I need the formulation.

Take any boolean function. Any class of tests. Partition, $\forall \varepsilon$ exists k, partition into k, such that in almost all partitions you can't do better than constant function in predicting it.

Fix set of vertices V of set n. Let U be edges in complete graph on V. (Interesting if U is not the complete graph, ex. U is the edges in d-regular expander on V.)

Let τ be the set of cuts defined by $A, B \subseteq V$. WLOG $A \cap B = \phi$.

Directed edge is $e = \{a, b\}$. T(e) = 1 iff $a \in A$ and $b \in B$. Apply this chain to this class of tests.

What does F_KT look like? You have a bunch of sets A, B and you're defined by which sets they go between. Or think of it as: for every test you have 3 regions: those in A, B, neither. When you look at all of k things, split vertices into 3^k regions. A little overkill, but all the tests, can split into 3^k . Knowing region of endpoints tells you which subset of tests are true.

This looks not quite quivalent. Here test sets are combinatorial rectangles. Sz: partition of left and right such that every rectangle works. That's a more structured version that what you want? One gives you the other?

If you use this kind of boosting, you don't get much better than this. If you use other types, you get more structured.

Plugging into the generic regularity lemma, $|E| \geq \delta\binom{n}{2}$. This says there exists $\mu = G(T_1, \ldots, T_k)$, where $k = O\left(\frac{1}{\varepsilon^2 \delta^2}\right)$ good predictor for any cut in the graph. This is the weak regularity of Kannan-Frieze. For Szemeredi we need the stronger boosting lemma.

Use the T's to divide vertices into 3^k subsets such that μ is a constant on every pair of subsets. Which sets you're in dictate which is true.

 3^k . If you want to know how many edges between A and B, this is

$$\frac{E_G(A,B)}{|E_G|} \approx_{\varepsilon} \sum_{i,j} \mu_{ij} \frac{|A \cap A_i||B \cap B_i|}{|V|^2}.$$

Constantly much information. You can't recover original set of edges, but can approximate any cut by knowing how big the intersections are relatively.

If we do this with G a subset of an expander, we get the same thing except

$$\frac{E_G(A,B)}{|E_{G'}|} =$$

Use expander mixing, with error term of expander mixing, get same thing.