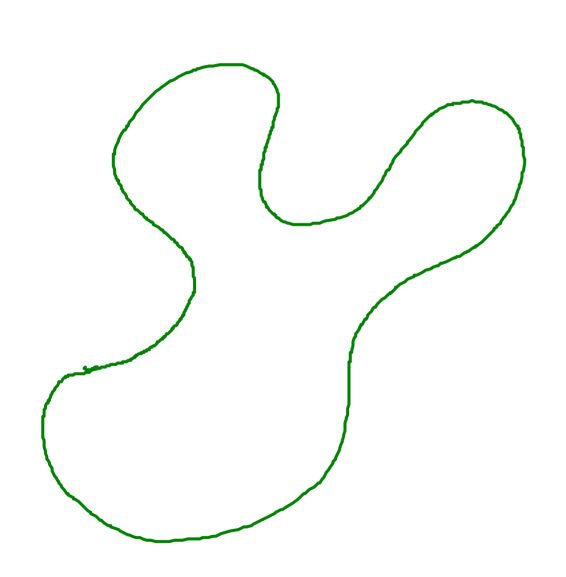
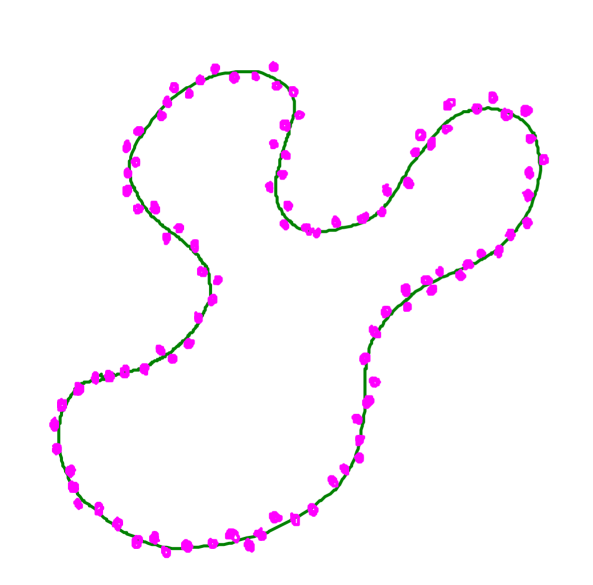
Niyogi-Smale-Weinberger Theorem

& its relatives

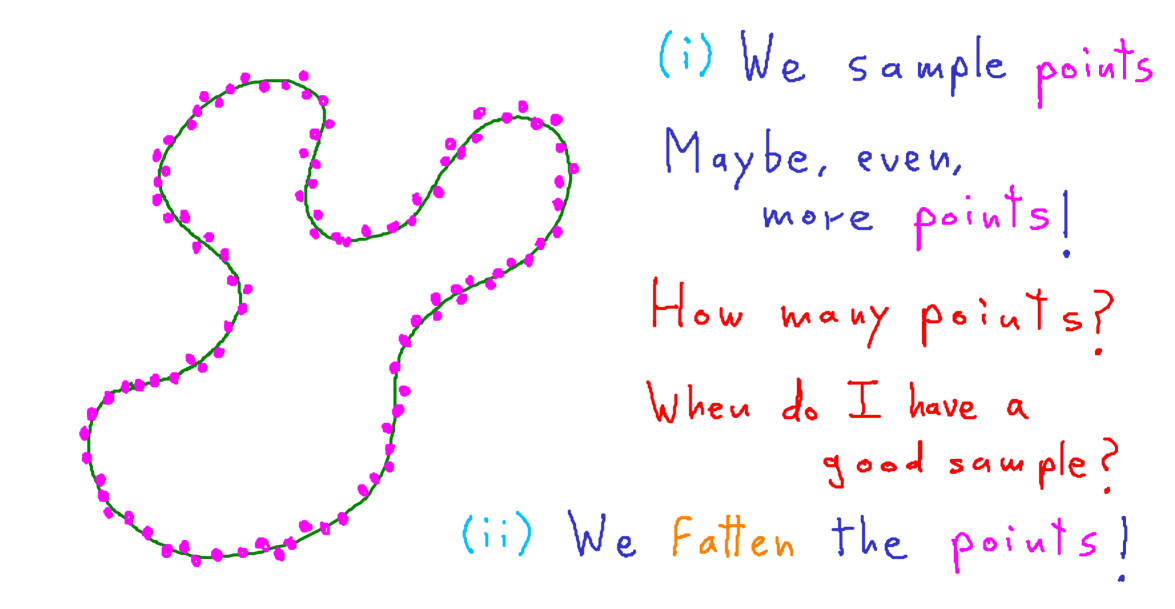
Josué TONELLI-CUETO

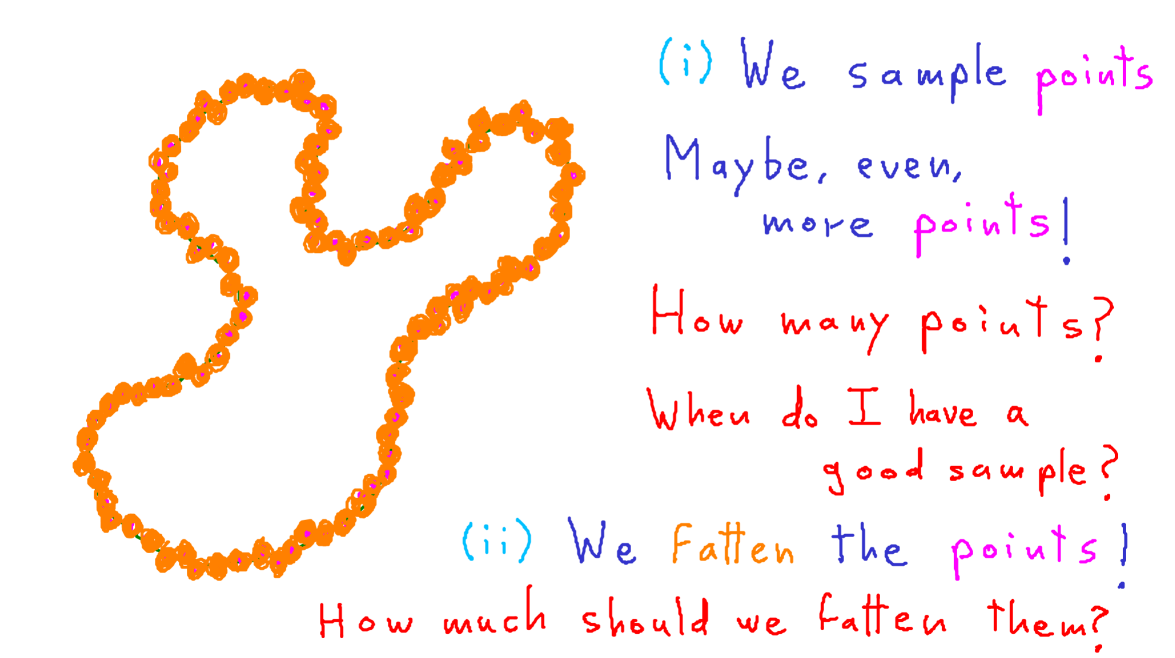


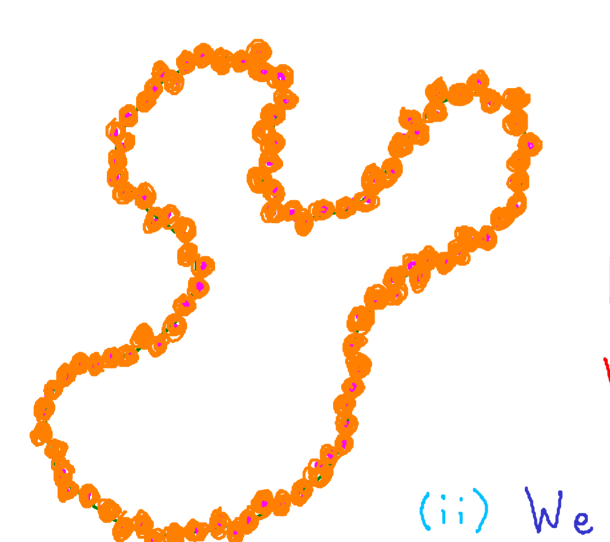
(i) We sample points



(i) We sample points Maybe, even,
more points! How many points? When do I have a good sample?







(i) We sample points Maybe, even, more points!

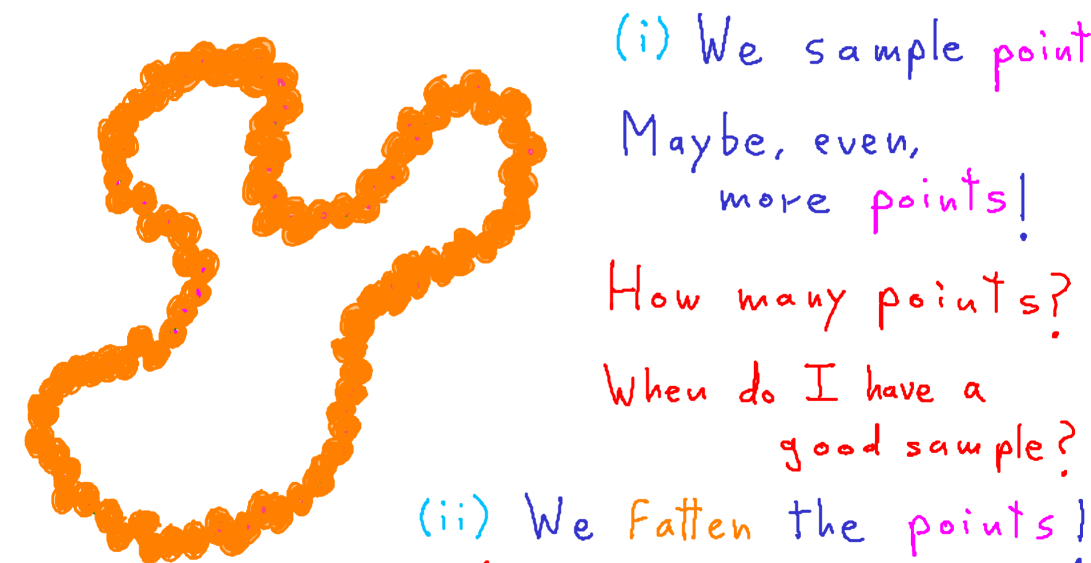
How many points?

When do I have a good sample?

(ii) We Fatten the points!

How much should we fatten them?

Maybe more!



(i) We sample points Maybe, even,
more points! How many points? When do I have a good sample?

How much should we fatten them? Maybe more! More?

FORMAL QUESTION:

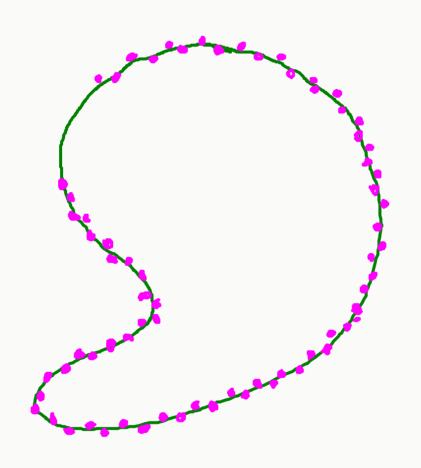
Given compact X CRM, a finite set $S \subseteq \mathbb{R}^m$ and $\epsilon > 0$, under which conditions do X and $B(S, \varepsilon) := \{x \in \mathbb{R}^m \mid dist(x, S) \leq \varepsilon \}$ "have the same topology"? (i.e. are of the same homotopy type?)

The "topology" of B(S, E) is that of the Zech complex of S and E and it can be computed... See other tutorials for more!

Smale what the NSW theorem is about Weinberger Niyogi How good is the sample? 1. Hausdorff distance 2 ingredients: 2. Reach (a.K.a. local Feature size)

Is the sample good enough?

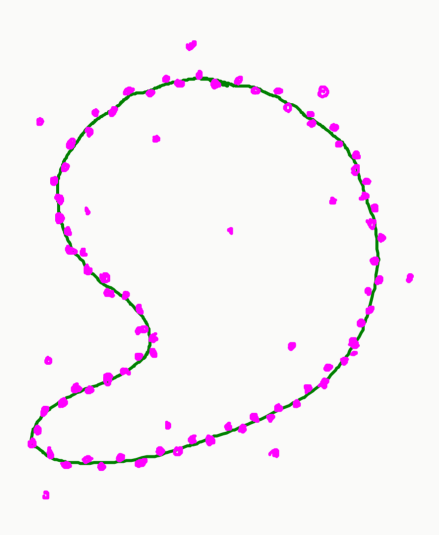
1st Ingredient: Haus dorff distance



Is this S a good sample of XP

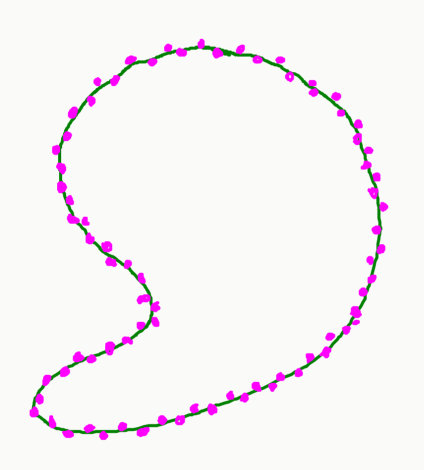
Nol Some points of X are too Far from 5

1 st Ingredient: Haus dorff distance



Is this S a good sample of X? Not Some points of S are too far from X

1 st Ingredient: Haus dorff distance



Is this S a 'good' sample of X? Maybe? Every point of X is 'near' S & every point of S is 'near' X

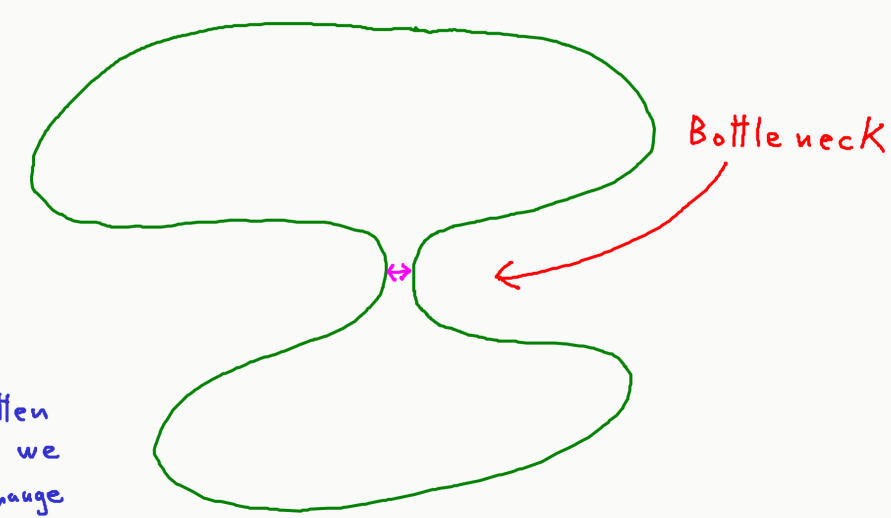
1 st Ingredient: Hans dorff distance

THM. disty is a metric on the set of non-empty compact subsets of IRM

dist H captures our intuitive notion of good sumple:

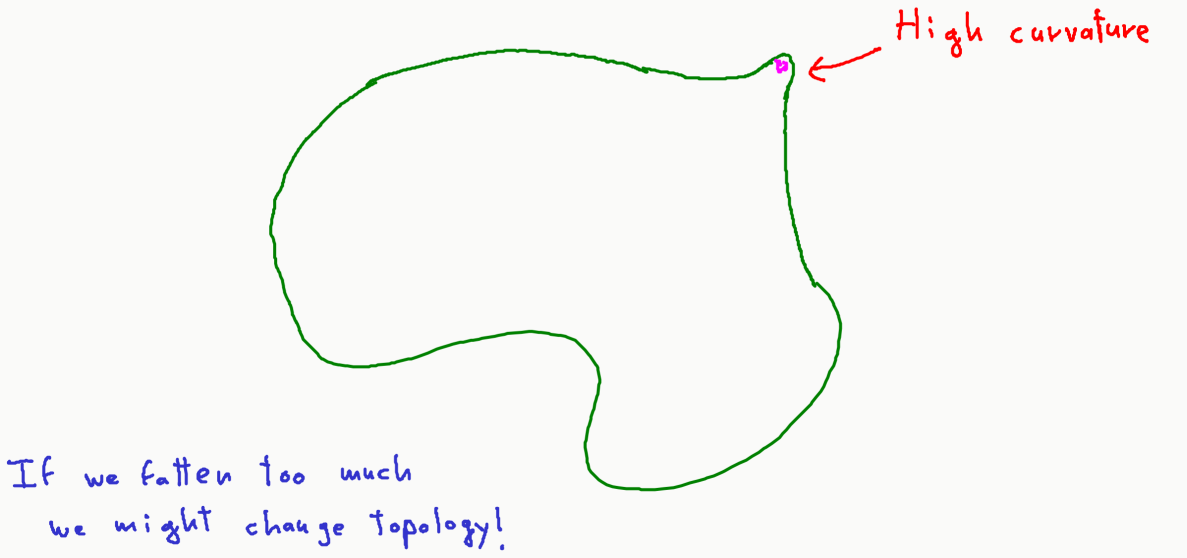
dist H (5, X) small (5) S is a good sumple of X

What can go wrong when we fatten the sample S of XP



If we fatten
too much we
might change
topology!

What can go wrong when we fatten the sample S of XP

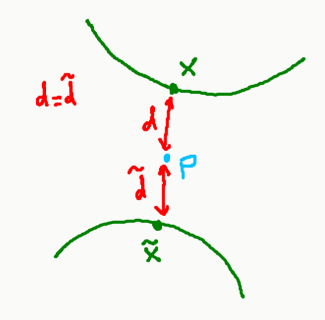


What can go wrong when we fatten the sample S of XP

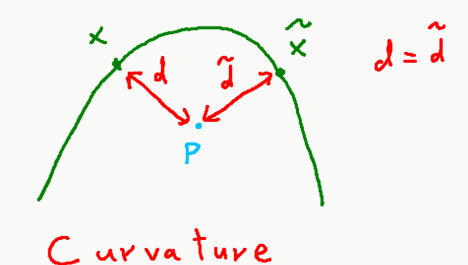
Medial axis:

$$\Delta_{X} := \{ P \in \mathbb{R}^{m} | \exists x, \hat{x} \in X : x \neq \hat{x}, dist(P,X) = dist(P,\hat{x}) \}$$

More than one nearest point in X



Bottleneck

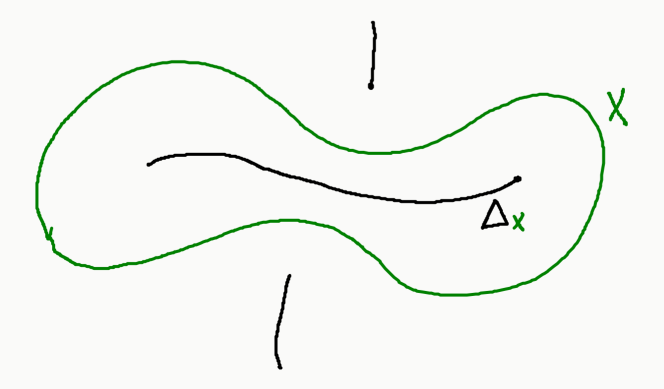


What can go wrong when we fatten the sample S of XP

Medial axis:

$$\Delta_{\chi} := \left\{ P \in \mathbb{R}^m | \exists x, \hat{x} \in \chi : x \neq \hat{x}, dist(P,\chi) = dist(P,\hat{x}) \right\}$$

More than one nearest point in X



What can go wrong when we fatten the sample S of X?

Medial axis:

$$\triangle_{\chi} := \left\{ p \in \mathbb{R}^m | \exists x, \hat{x} \in \chi : x \neq \hat{x}, dist(p, \chi) = dist(p, \hat{x}) \right\}$$

More than one nearest point in X

Reach:

$$\gamma(\chi) := \inf_{x \in X} dist(x, \Delta_{\chi})$$

T(X) measures how hard is to sample X:

T(X) 'small' (=) X is 'hard' to sample

NSW theorem: compact set X finite set 5 e > 0If (i) $dist_H(S, X) \leq (\sqrt{9} - \sqrt{8}) \gamma(X)$ and (iii) $\frac{d_{s}t_{H}(S,X)+r(X)-\sqrt{d_{s}t_{H}(S,X)^{2}+r(X)^{2}-6d_{s}t_{H}(S,X)r(X)}}{2}$ < & < $dist_{H}(S,X)+r(X)+\sqrt{dist_{H}(S,X)^{2}+r(X)^{2}-6dist_{H}(S,X)}r(X)$

Then

B(5, E) and X are of the same homotopy type.

1 S W theorem (Easier to read version) compact set X finite set 5 e > 0IF $3 \operatorname{dist}_{H}(5,X) < \varepsilon < \frac{1}{2} \gamma(X)$

Then B(5, E) and X are of the same homotopy type.

```
Can we do more?
YES...
```

- + Weak Reach (Chazal, Lieutier; 2005)
- + Balls of different radii (Chazal, Lientier; 2007) (Han; 2019) (Eckhardt; 2020)
- + Vietoris-Rips complex (Attali, Lieutier, Salinas; 2012)
- + Ellipsoids (Kališnik, Lešnik; 2020+)
 - ... and much more!



D. Attali, A. Lieutier, and D. Salinas.

Vietoris-Rips complexes also provide topologically correct reconstructions of sampled shapes.

Comput. Geom., 46(4):448-465, 2013.



E Chazal and A Lieutier.

Weak feature size and persistant homology: computing homology of solids in \mathbb{R}^n from noisy data samples.

In Computational geometry (SCG'05), pages 255–262. ACM, New York, 2005.



F. Chazal and A. Lieutier.

Smooth manifold reconstruction from noisy and non-uniform approximation with guarantees.

Comput. Geom., 40(2):156-170, 2008.



A. Eckhardt.

An Adaptive Algorithm for Computing the Homology of Semialgebraic Sets.

Master's thesis, Technische Universität Berlin, 2020.



I. Han.

An Adaptive Grid Algorithm for Computing the Homology Group of Semialgebraic Set.

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S. Kališnik and D. Lešnik.

Finding the homology of manifolds using ellipsoids, 2020. arXiv:2006.09194.



P. Niyogi, S. Smale, and S. Weinberger.

Finding the homology of submanifolds with high confidence from random samples.

Discrete Comput. Geom., 39(1-3):419-441, 2008.