## Agnesi Function Based Ionic Particle System

### This system presents the following properties:

- Function based
- Emergent behavior from the math properties themselves
- Conceptually applicable to physical analog replicas
- Simple (without excesive mathematical complexity)
- for a conventional digital computing system

## Particle Representation Using Agnesi Function

$$f(x) = \frac{1}{x^2 + 1}$$



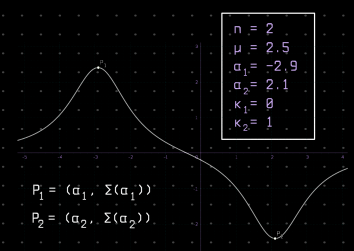
Horizontal Asymptote

$$\lim_{x\to\infty} \frac{1}{x^2+1} = 0 \qquad f(x)>0 \quad \forall \quad x \in \mathbb{R}$$

Particle

$$\rho_{i}(x) = \frac{(-1)^{K_{i}} \mu}{(x-\alpha_{i})^{2}+1} \quad \alpha_{i}, \mu \in \mathbb{R}, \kappa_{i} \in \mathbb{N}$$

### Example field



I chose this function because it' energy that influences the entire field around it. There's no distance where the other particles can't sense that energy. That's caused by the horizontal asymptote of the function

Original	Variable Particle Position				Variable Mass Coefficent			
1		• •1 •				. µ		
$\times^2+1$	<b>→</b>	(x-a)2+	-1	٠	7	(x-a)	2+1	

Variable Particle

$$\frac{1}{1+\alpha} \Rightarrow \frac{(-1)^{K} \mu}{(x-\alpha)^{2}+1}$$

The mass coefficient (μ) is a global parameter The particle charge  $((-1)^{\prime}$ allows the mass to be

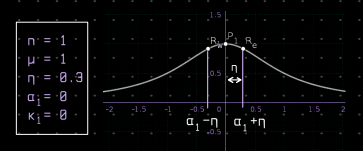
$$\sum_{i=1}^{n} p_{i}(x)$$

The field represents the sum of each particle function

The field is the function. The energy is the value of the field at a

### Interactions

In order to move, the only information the particle needs to know is the energy at certain short distance in each side of its position. I named that distance η

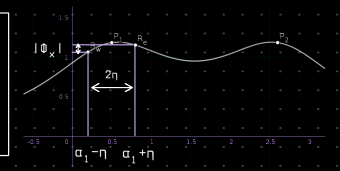


In R¹ we got just 2 reference values per particle: α,-η, and α,+η

### Force

The force is the difference between the energy in the right reference value and the left one

$$\Phi_{\times} = \sum_{i} (\alpha_{i} + \eta_{i}) - \sum_{i} (\alpha_{i} - \eta_{i})$$



When there's more energy at the left side, the force is negative. If it's at the right side, the force is

positive. That force value will

increase as the particle moves towards the energy source

## Translating to $\mathbb{R}^2$ (2D)

Original

$$f(x) = \frac{1}{x^2 + 1}$$

$$g(x;y) = f(\sqrt{(x^2+y^2)})$$

$$\frac{1}{(\sqrt{(x^2+y^2)})^2+1} = \frac{1}{x^2+y^2}$$

(α<sub>i</sub>-η, β<sub>i</sub>)•

Particle

$$\rho_{i}(x) = \frac{(-1)^{\kappa_{i}} \mu}{(x-\alpha_{i})^{2} + (y-\beta_{i}) + 1} \alpha_{i}, \mu \in \mathbb{R}, \kappa_{i} \in \mathbb{N}$$

Field

n = 2

μ = 1

 $\kappa_1 = 0$ 

 $\kappa_2 = 0$ 

 $\eta = 0.3$  $a_1 = 0.5$  $a_{2} = 2.6$ 

$$\Sigma(x,y) = \sum_{i=1}^{n} \rho_{i}(x,y)$$

Forces



$$\Phi_{y} = \Sigma(\alpha_{i}, \beta_{i} + \eta) - \Sigma(\alpha_{i}, \beta_{i} - \eta)$$

# $(\alpha_i, \beta_i - \eta)$

With this parameters, now the particles will follow the lowest energy path. In consequence the particles of opposite charge will attract each other and the ones with same charge are going to repel each other. Everything is going to converge into energetically balanced structures

## Position Update

Now with everything in place. The next step is to update the positions using the force values. To adjust the speed and direction of the interactions, i'm going to set a global force multiplier value named  $\Omega$ 

$$\alpha_{i} = \alpha_{i} + \Omega \Phi_{\times}$$

Ω>0 ⇒ Go towards energy

$$\beta_i = \beta_i + \Omega \Phi_y$$

Ω<0 ⇒ Go against energy