D. Excitable Media

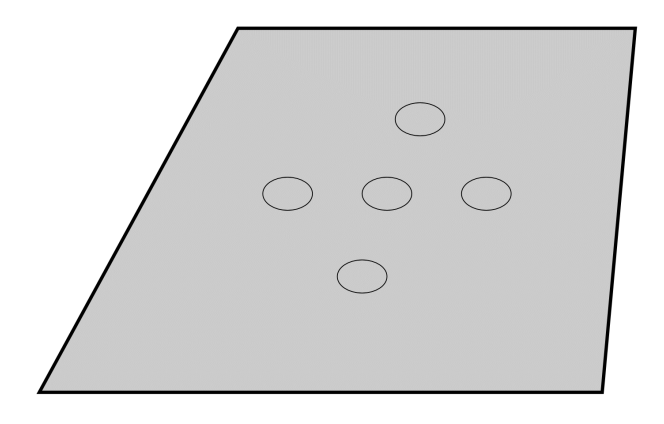
Examples of Excitable Media

- Slime mold amoebas
- Cardiac tissue (& other muscle tissue)
- Cortical tissue
- Certain chemical systems (e.g., BZ reaction)
- Hodgepodge machine

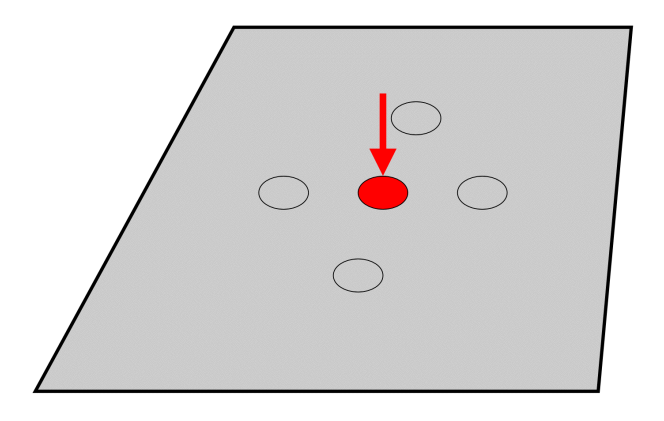
Characteristics of Excitable Media

- Local spread of excitation
 - for signal propagation
- Refractory period
 - for unidirectional propagation
- Decay of signal
 - avoid saturation of medium

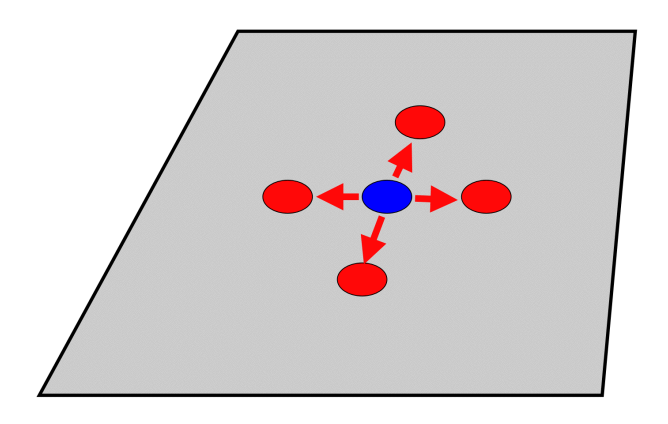
Behavior of Excitable Media



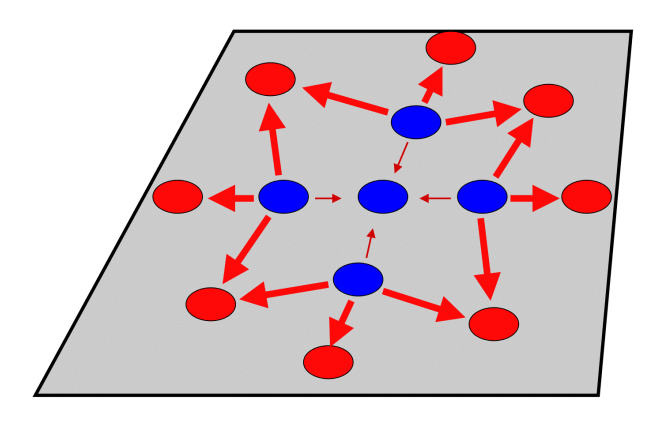
Stimulation



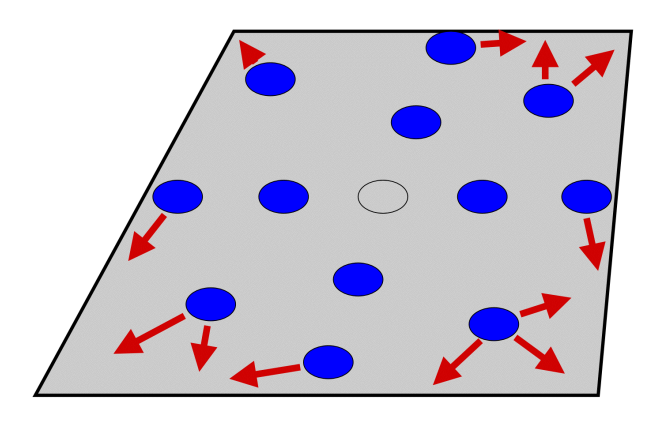
Relay (Spreading Excitation)



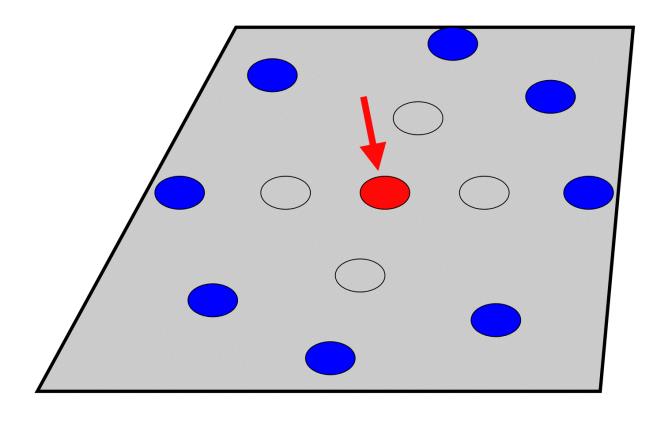
Continued Spreading



Recovery



Restimulation



Circular & Spiral Waves Observed in:

- Slime mold aggregation
- Chemical systems (e.g., BZ reaction)
- Neural tissue
- Retina of the eye
- Heart muscle
- Intracellular calcium flows
- Mitochondrial activity in oocytes

Cause of Concentric Circular Waves

- Excitability is not enough
- But at certain developmental stages, cells can operate as pacemakers
- When stimulated by cAMP, they begin emitting regular pulses of cAMP

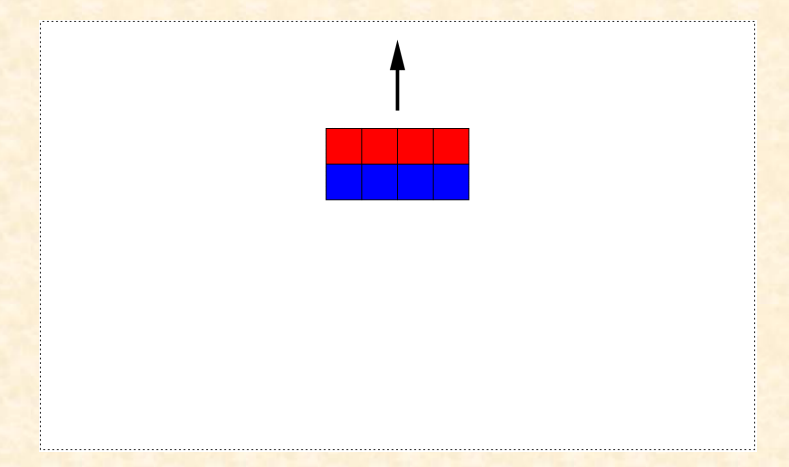
Spiral Waves

- Persistence & propagation of spiral waves explained analytically (Tyson & Murray, 1989)
- Rotate around a small core of non-excitable cells
- Propagate at higher frequency than circular
- Therefore they dominate circular in collisions
- But how do the spirals form initially?

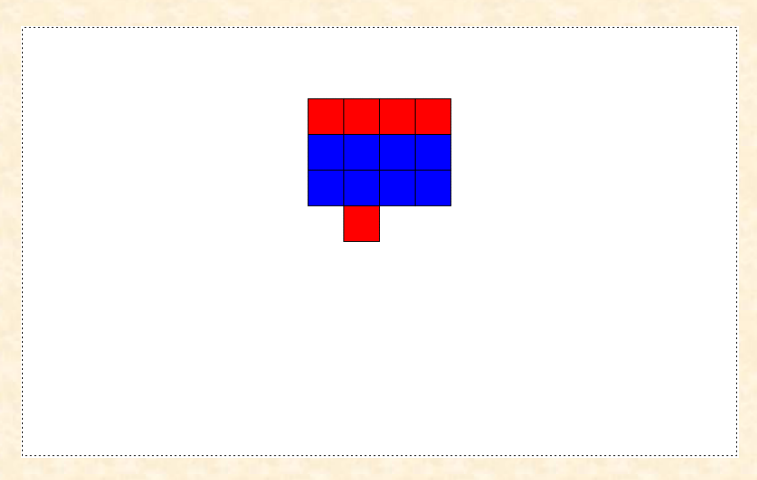
Some Explanations of Spiral Formation

- "the origin of spiral waves remains obscure" (1997)
- Traveling wave meets obstacle and is broken
- Desynchronization of cells in their developmental path
- Random pulse behind advancing wave front

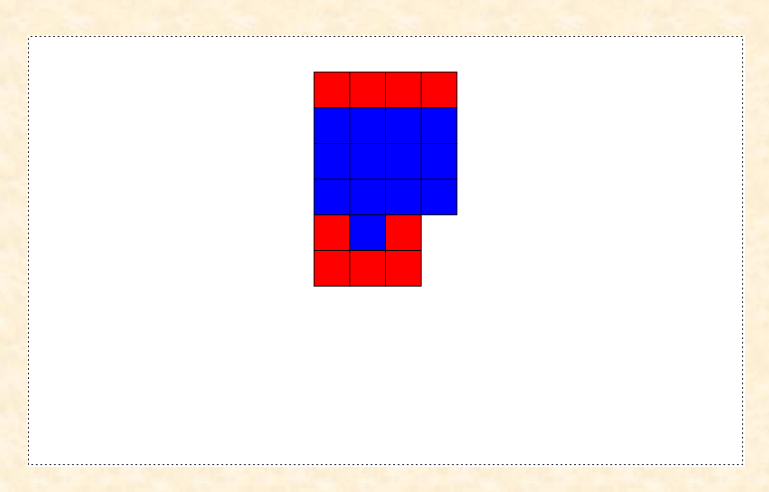
Step 0: Passing Wave Front

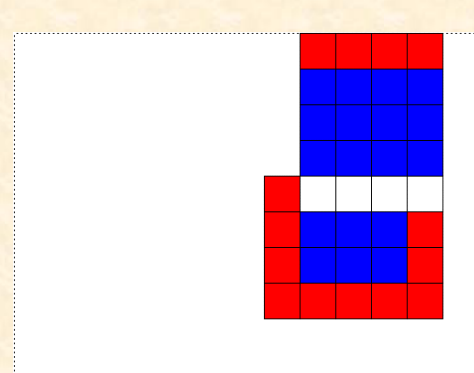


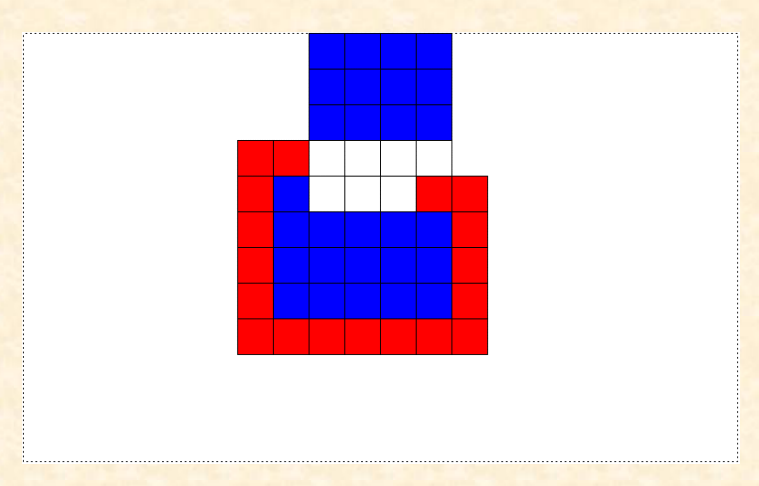
Step 1: Random Excitation

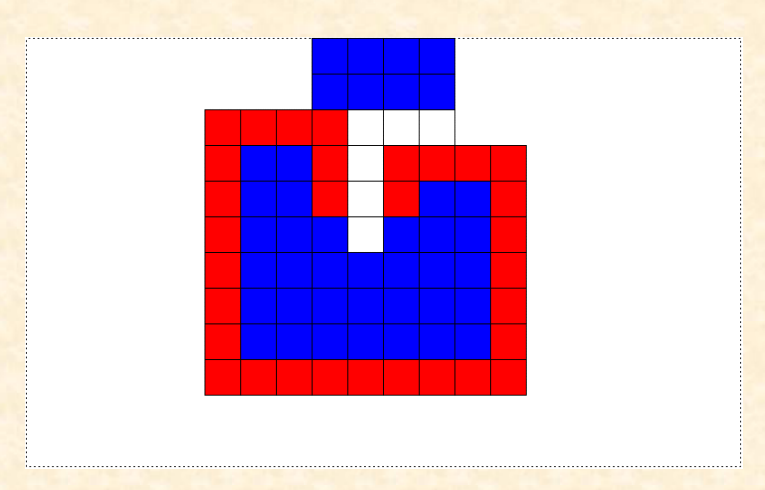


Step 2: Beginning of Spiral

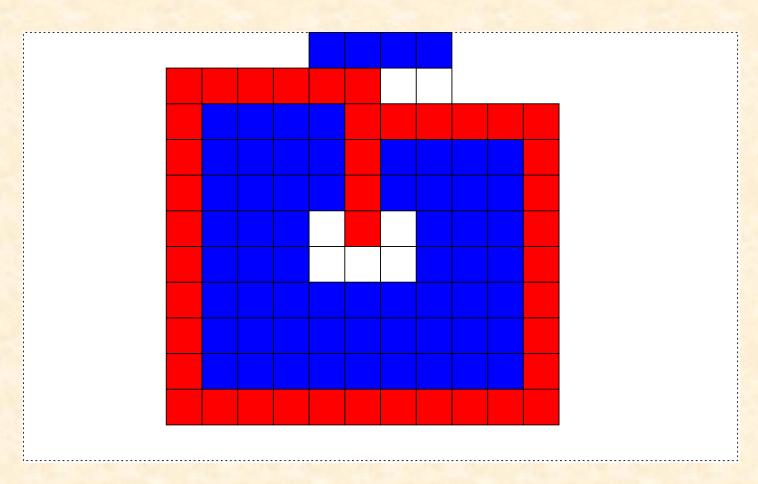




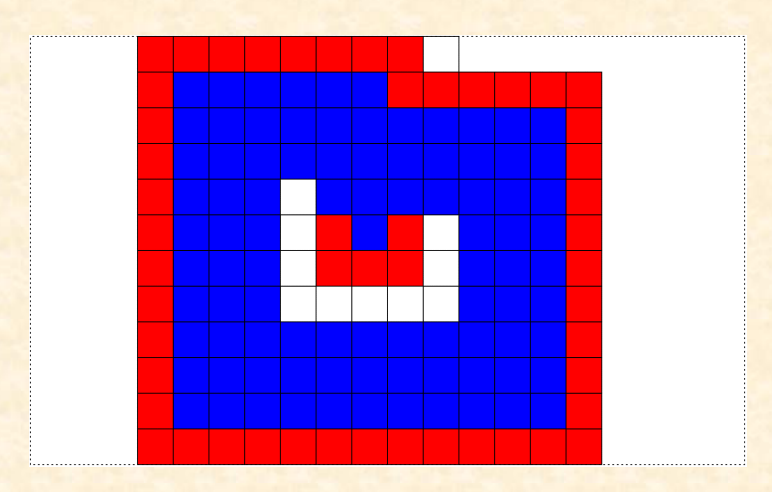


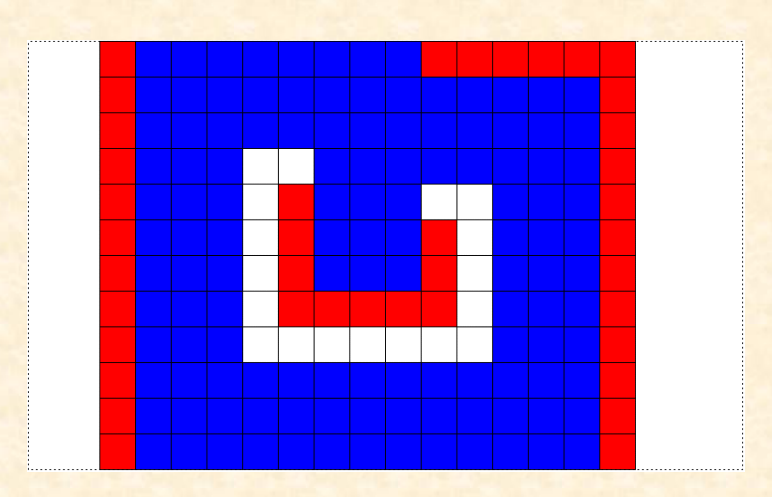


Step 6: Rejoining & Reinitiation

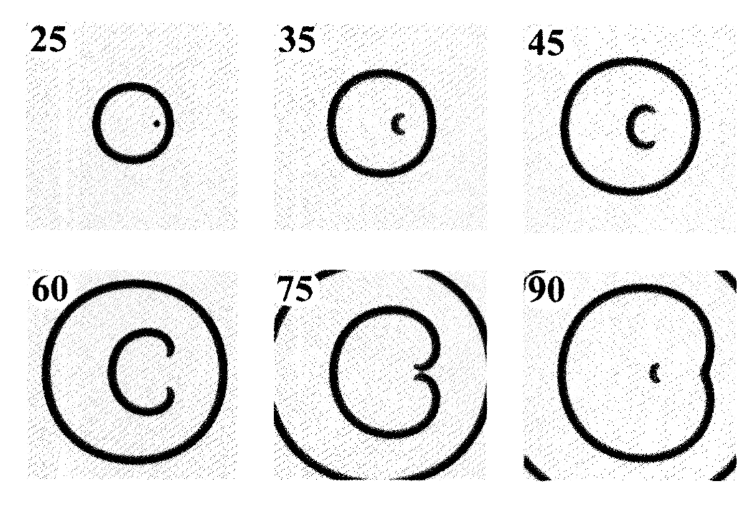


Step 7: Beginning of New Spiral





Formation of Double Spiral



from Pálsson & Cox (1996)

NetLogo Simulation Of Spiral Formation

- Amoebas are immobile at timescale of wave movement
- A fraction of patches are inert (grey)
- A fraction of patches has initial concentration of cAMP
- At each time step:
 - chemical diffuses
 - each patch responds to local concentration

Response of Patch

```
if patch is not refractory (brown) then
  if local chemical > threshold then
    set refractory period
    produce pulse of chemical (red)
else
```

decrement refractory period degrade chemical in local area

Demonstration of NetLogo Simulation of Spiral Formation

Run SlimeSpiral.nlogo

Demonstration of NetLogo Simulation of Spiral Formation (a closer look)

Run SlimeSpiralBig.nlogo

Observations

- Excitable media can support circular and spiral waves
- Spiral formation can be triggered in a variety of ways
- All seem to involve inhomogeneities (broken symmetries):
 - in space
 - in time
 - in activity
- Amplification of random fluctuations
- Circles & spirals are to be expected

NetLogo Simulation of Streaming Aggregation

- 1. chemical diffuses
- 2. **if** cell is refractory (yellow)
- 3. then chemical degrades
- 4. **else** (it's excitable, colored white)
 - 1. **if** chemical > movement threshold **then** take step up chemical gradient
 - 2. **else if** chemical > relay threshold **then** produce more chemical (red) become refractory
 - 3. **else** wait

Demonstration of NetLogo Simulation of Streaming

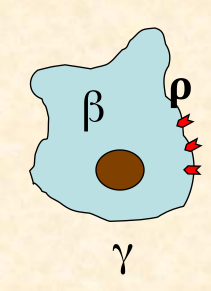
Run SlimeStream.nlogo

Model for Dicty Signalling

Variables (functions of x, y, t):

 β = intracellular concentration of cAMP

γ = extracellular concentration of cAMP



 ρ = fraction of receptors in active (sensitive) state

Equations

$$\frac{d\beta(x,y,t)}{dt} = s\Phi(\rho,\gamma) \qquad -\beta k_{\rm i} \qquad -\beta k_{\rm t} \qquad [1]$$

 $\begin{array}{ll} \text{Rate of change in} \\ \text{intracellular [cAMP]} = \begin{array}{ll} \text{Production} \\ \text{of cAMP} \end{array} & - \begin{array}{ll} \text{Intracellular} \\ \text{hydrolysis} \end{array} - \begin{array}{ll} \text{Secretion} \\ \text{of cAMP} \end{array}$

$$\frac{d\gamma(x,y,t)}{dt} = \frac{k_t}{h}\beta \qquad -k_e\gamma \qquad +D\nabla^2\gamma \quad [2]$$

$$\frac{d\rho(x,y,t)}{dt} = f_2(\gamma)(1-\rho) - f_1(\gamma)\rho$$
 [3]

 $\frac{\text{Rate of change in fraction of active receptor}}{\text{tion of active receptor}} = \frac{\text{Dephospho-}}{\text{rylation of receptor}} - \frac{\text{Phosphorylation}}{\text{of receptor}}$

Positive Feedback Loop

- Extracellular cAMP increases
 (γ increases)
- → Rate of synthesis of intracellular cAMP increases

(Ф increases)

- ⇒ Intracellular cAMP increases (β increases)
- ⇒ Rate of secretion of cAMP increases
- (⇒ Extracellular cAMP increases)

Negative Feedback Loop

- Extracellular cAMP increases (γ increases)
- \Rightarrow cAMP receptors desensitize (f_1 increases, f_2 decreases, ρ decreases)
- → Rate of synthesis of intracellular cAMP decreases

(Φ decreases)

- → Intracellular cAMP decreases
 (β decreases)
- ⇒ Rate of secretion of cAMP decreases
- ⇒ Extracellular cAMP decreases
 (γ decreases)

Dynamics of Model

- Unperturbed
 - ⇒ cAMP concentration reaches steady state
- Small perturbation in extracellular cAMP
 - ⇒ returns to steady state
- Perturbation > threshold ⇒
 - large transient in cAMP, and then return to steady state
 - or oscillation (depending on model parameters)

Typical Equations for Excitable Medium (ignoring diffusion)

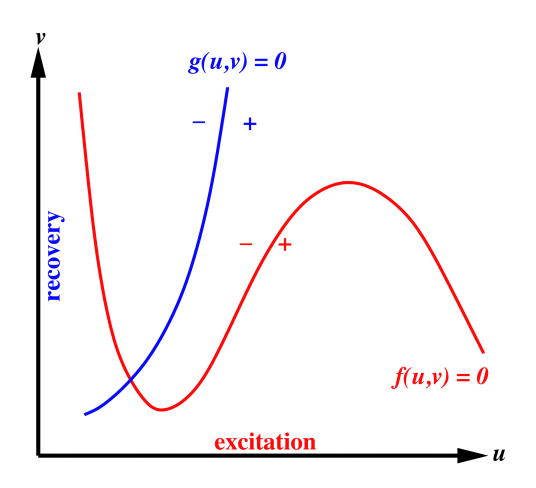
• Excitation variable:

$$\dot{u} = f(u,v)$$

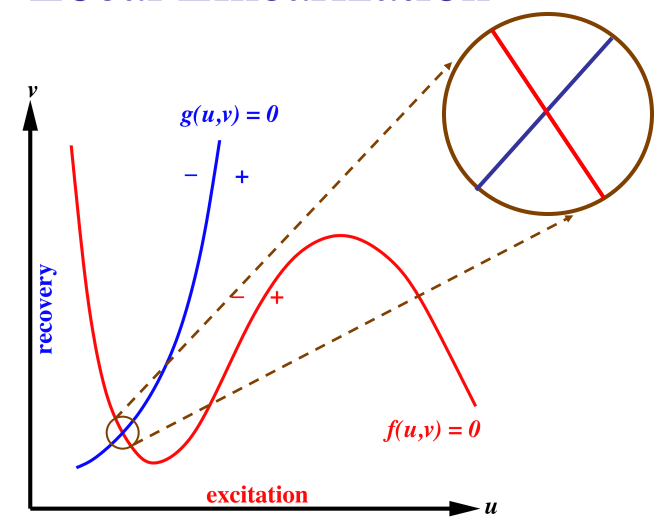
• Recovery variable:

$$\dot{v} = g(u,v)$$

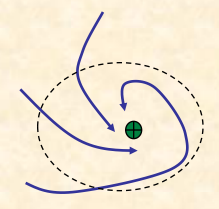
Nullclines

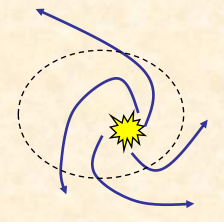


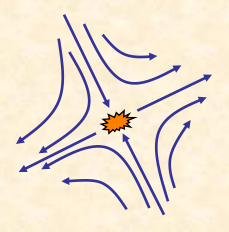
Local Linearization



Fixed Points & Eigenvalues







stable fixed point

real parts of eigenvalues are negative

unstable fixed point

real parts of eigenvalues are positive

saddle point

one positive real & one negative real eigenvalue

Neural Impulse Propagation

$$C\frac{dv}{dt} = I - g_{Na}m^3h(V - V_{Na}) - g_Kn^4(V - V_K) - g_L(V - V_L)$$

$$\frac{dm}{dt} = a_m(V)(1 - m) - b_m(V)m$$

$$\frac{dh}{dt} = a_h(V)(1 - h) - b_h(V)h$$

$$\frac{dn}{dt} = a_n(V)(1 - n) - b_n(V)n$$

$$a_m(V) = .1(V + 40)/(1 - \exp(-(V + 40)/10))$$

$$b_m(V) = 4\exp(-(V + 65)/18)$$

$$a_h(V) = .07\exp(-(V + 65)/20)$$

$$b_h(V) = 1/(1 + \exp(-(V + 35)/10))$$

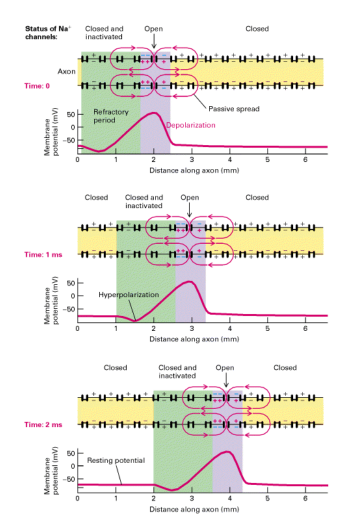
$$a_n(V) = .01(V + 55)/(1 - \exp(-(V + 55)/10))$$

$$b_n(V) = .125\exp(-(V + 65)/80)$$
Time:1 ms

Time:1 ms

Time:1 ms





FitzHugh-Nagumo Model

- A simplified model of action potential generation in neurons
- The neuronal membrane is an excitable medium
- B is the input bias:

$$\dot{u} = u - \frac{u^3}{3} - v + B$$

$$\dot{v} = \varepsilon (b_0 + b_1 u - v)$$

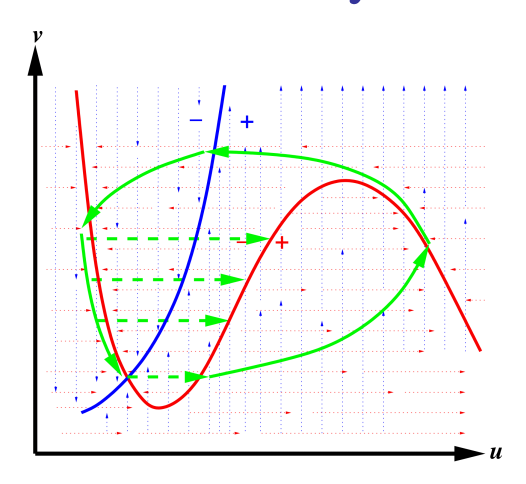
Nullclines

- u nullcline: $v = u \frac{u^3}{3} + B$
- v nullcline: $v = b_0 + b_1 u$

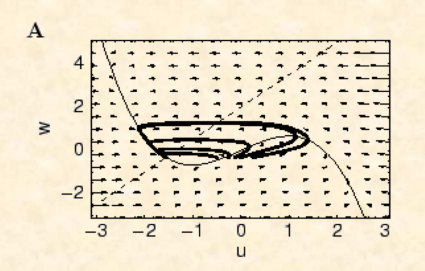
NetLogo Simulation of Excitable Medium in 2D Phase Space

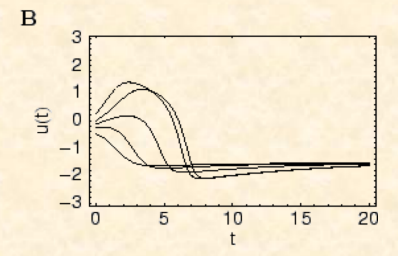
(EM-Phase-Plane.nlogo)

Elevated Thresholds During Recovery



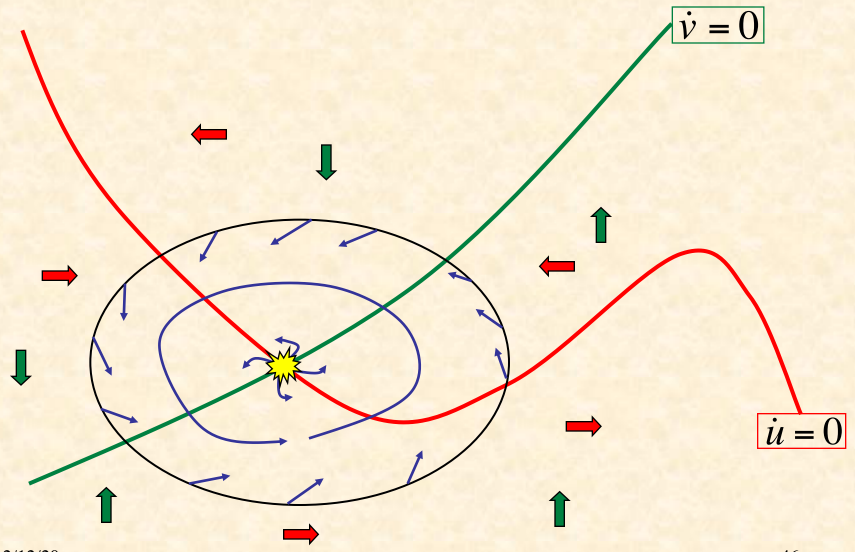
Type II Model



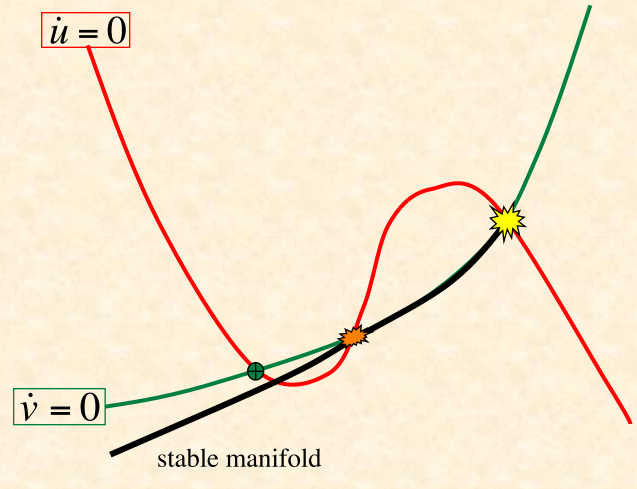


- Soft threshold with critical regime
- Bias can destabilize fixed point

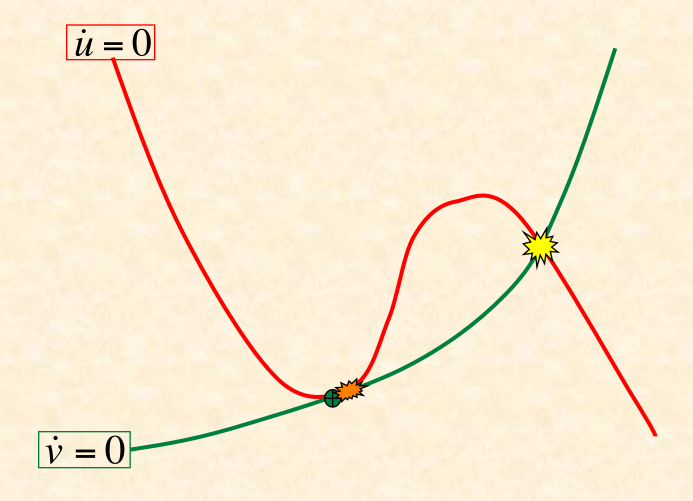
Poincaré-Bendixson Theorem



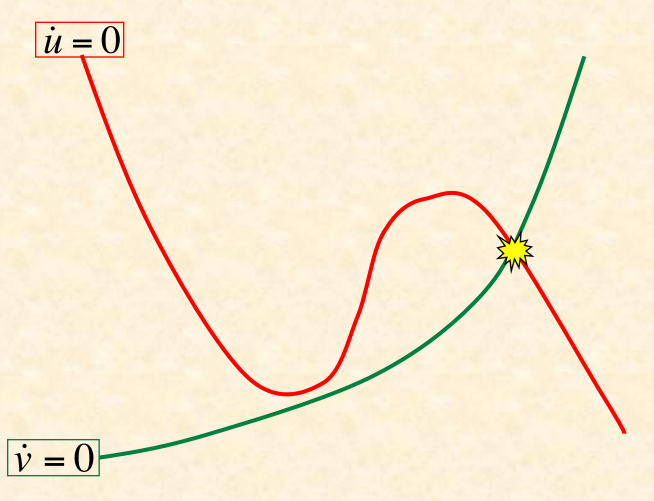
Type I Model



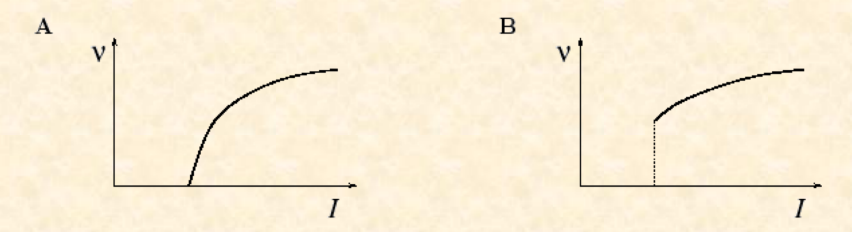
Type I Model (Elevated Bias)



Type I Model (Elevated Bias 2)



Type I vs. Type II



- Continuous vs. threshold behavior of frequency
- Slow-spiking vs. fast-spiking neurons

fig. < Gerstner & Kistler



Additional Bibliography

- 1. Kessin, R. H. Dictyostelium: Evolution, Cell Biology, and the Development of Multicellularity. Cambridge, 2001.
- 2. Gerhardt, M., Schuster, H., & Tyson, J. J. "A Cellular Automaton Model of Excitable Media Including Curvature and Dispersion," *Science* **247** (1990): 1563-6.
- 3. Tyson, J. J., & Keener, J. P. "Singular Perturbation Theory of Traveling Waves in Excitable Media (A Review)," *Physica D* **32** (1988): 327-61.
- 4. Camazine, S., Deneubourg, J.-L., Franks, N. R., Sneyd, J., Theraulaz, G., & Bonabeau, E. *Self-Organization in Biological Systems*. Princeton, 2001.
- 5. Pálsson, E., & Cox, E. C. "Origin and Evolution of Circular Waves and Spiral in *Dictyostelium discoideum* Territories," *Proc. Natl. Acad. Sci. USA*: **93** (1996): 1151-5.
- 6. Solé, R., & Goodwin, B. Signs of Life: How Complexity Pervades Biology. Basic Books, 2000.