

Enzyme kinetics

(called substrate)

Enzymes are catalysts - they help convert other molecules into products but are not used up in the reaction themselves.

Consider chemicals A and B reacting on collision to form chemical C, with a rate k:



this depends on the molecule shapes and sizes and the temperature.

Then we can write

$$\frac{dC}{dt} = kAB$$

More on this
on problem sheet
1 question 2

Get students to think why this is true - if you double the number of A you would expect the rate of reaction to double

N/Ae this means that

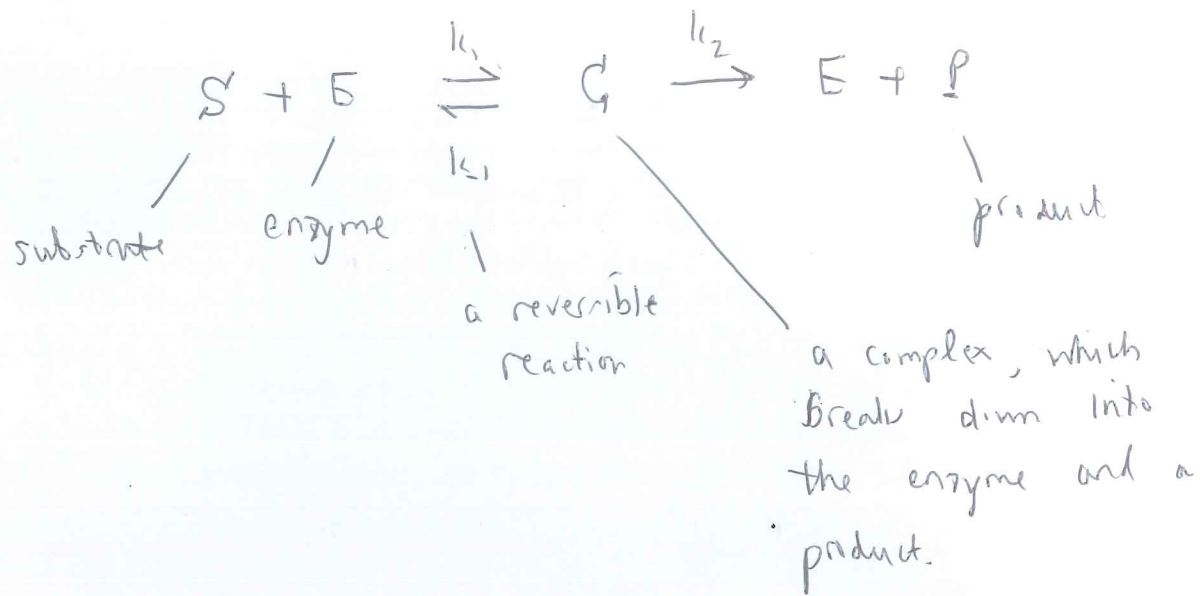


$$\text{is } \frac{dC}{dt} = kA^2B$$

This is called the law of mass action

Assumes that the mixture is well stirred (no spatial dependence).

An enzyme reaction looks like this:



The overall reaction is $S \rightarrow P$. This looks like a simple reaction, but we know there are intermediate steps. You could just model the single reaction and capture all the intermediate steps with this overall reaction. This is the Mil equation:

$$r = \frac{k_2 S^n}{K_m + S^n}$$

some constants

But better to use the law of mass action:

$$\begin{aligned}
 \frac{dS}{dt} &= k_{-1} S - k_1 S E & \text{①} \\
 \frac{dE}{dt} &= (k_{-1} + k_2) C - k_1 S E & \text{②} \\
 \frac{dC}{dt} &= k_1 S E - (k_2 + k_{-1}) C & \text{③} \\
 \frac{dP}{dt} &= k_2 C & \text{④}
 \end{aligned}$$

Can we simplify? Yes -

3.

I only appears in $\textcircled{4}$ so decouples

Add $\textcircled{2} + \textcircled{3}$ to ~~eliminate~~ see that $E + C = \text{constant}$

$$\begin{array}{l} \text{Initial} \\ \text{value of } E \end{array} \quad \underline{\hspace{10em}} \quad = E_0$$

(since $C = 0$ initially)

This reduces the system to two ODEs:

$$\frac{dS}{dt} = k_1 C - k_1 S (E_0 - C)$$

$$\frac{dC}{dt} = k_1 S (E_0 - C) - (k_{-1} + k_1) C$$

Which can be solved subject to suitable initial conditions $S = S_0, C = 0$ at $t = 0$.

Non-dimensionalize to analyse the system:

$$S = S_0 s, \quad C = E_0 c, \quad t = \frac{t'}{k_1 E_0}$$

gives

$$\frac{ds}{dt'} = -s + c(s + k' - \lambda)$$

$$k' = \frac{k_{-1} + k_2}{k_1 S_0}$$

$$\frac{dc}{dt'} = s - (s + k')c$$

$$\lambda = \frac{k_2}{k_1 S_0}$$

$$s(0) = 1, \quad c(0) = 0$$

$$\epsilon = \frac{E_0}{S_0} \ll 1$$

because we only need a bit of enzyme

$s \ll 1$ means we can neglect the time derivative in the c equation. This makes a quasistatic system: s evolve through a time derivative; c evolves through an algebraic equation.

More on this in problem sheet 1, question 1!

$\Rightarrow c = \frac{s}{s + k'}$

$$\frac{ds}{dt} = - \frac{\lambda s}{s + k'} \quad (*)$$

k' is known as the Michaelis-Menten law (and is for enzyme reactions).

What is the reaction rate?

This is $r \stackrel{\text{def}}{=} \frac{df}{dt} = - \frac{ds}{dt}$

(because it is minus the rate of depletion)

$= -s_0 E_0 k_1 \frac{ds}{dt}$ (non-dimensionalising)

$= \frac{k_2 E_0 s}{k + s}$ using (*)

where $k = \frac{k_{-1} + k_2}{k_1}$

is the Michaelis constant.

It's difficult to measure individual reaction rates experimentally. But we can measure the overall reaction rate r and concentrations. ~~But~~ so, if we look at the initial rate of reaction r_0 :

$$r_0 = \frac{k_2 E_0 S_0}{k + S_0} \Rightarrow \frac{1}{r_0} = \frac{1}{k_2 E_0} + \left(\frac{k}{k_2 E_0} \right) \frac{1}{S_0}$$

so $\frac{1}{r_0}$ is linear in $\frac{1}{S_0}$.

We can plot experimental data of $\frac{1}{r_0}$ versus $\frac{1}{S_0}$ and the gradient is $\frac{k}{k_2 E_0}$ and intercept is $\frac{1}{k_2 E_0}$ which are for different S_0 's.

allows us to extract k and $k_2 E_0$. These plots are called Lineweaver-Burk plots.

our quasi-steady approximation does not satisfy our initial conditions. Now $c = \frac{s}{s+k}$ ~~is~~ $c = 0$. This is because there is a rapid transient. We see this by rescaling $t' = \varepsilon \tau$ to give

$$\frac{ds}{dt} = \varepsilon (-s + c(s + k) - \lambda) = 0 \text{ to leading order} \Rightarrow s = s_0$$

$$\frac{dc}{dt} = s - (s + k)c$$

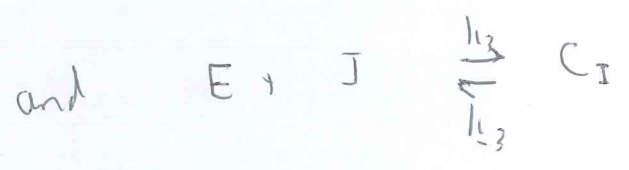
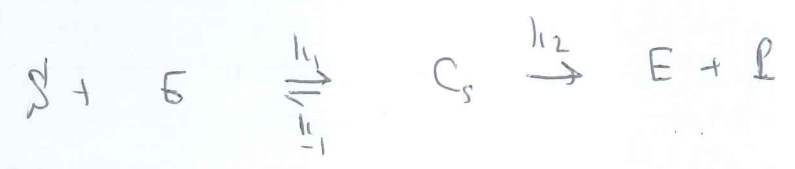
$$\Rightarrow \frac{dc}{dt} = s_0 - (s_0 + k)c$$

$$\Rightarrow c = \frac{1}{1+k} (1 - e^{-(1+k)t})$$

This takes us from $c=0$ at $t=0$ to the initial quasi-steady value.

Inhibitors are substances that inhibit the catalytic reaction of an enzyme ~~eg by blocking reaction sites~~

eg 1 Competitive inhibition - when the substrate can't bind if the inhibitor is bound to an enzyme

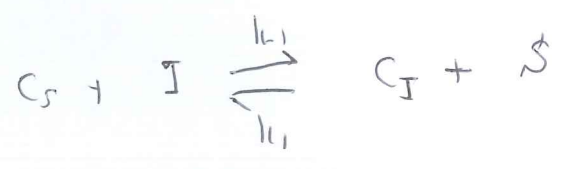


You can do law of mass action for these and analyse the reaction rate in a similar way to the previous case (see lecture notes)

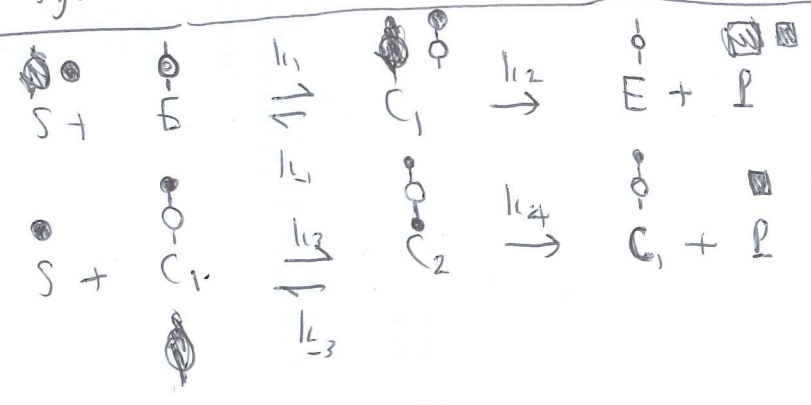
eg 2 allosteric inhibition

As well as the previous two reactions we can also have

the inhibitor binding to C_S to make a different product:



Cooperative systems - more than one binding site



(catalytic reaction to make P from S using E as a catalyst)
 (catalytic reaction to make P from S using the intermediate product C_1 as a catalyst too)

You can do the law of mass action analysis in this case and find that now

$$r = \frac{(k_2 k_2 + k_4 S') E_0 S}{k_1 k_2 + k_2 S' + S'^2}$$

$$K_1 = \frac{k_{-1} + k_2}{k_1}$$

$$K_2 = \frac{k_4 + k_3}{k_3}$$

Limiting

Special cases: if the rates of binding and reaction are equal at each site identical and independent at each site then $k_1 = 2k_3$ (because k_3 is a reaction binding S with C_1 and C_1 has because there are

Special case: if the rate of binding of the first substrate molecule is small but the rate of binding of the second molecule is large then $k_1 \rightarrow 0$, $k_3 \rightarrow \infty$ with k_1, k_3

Finite gives

$$r = \frac{k_4 E_0 S^2}{k_1 k_2 + S^2}$$

which is a Hill equation with exponent 2

Trans-membrane ion transport

Cells are bags of fluid water

The water contains dissolved salts: NaCl and KCl which dissolve into Na^+ , Cl^- , K^+ , ~~or~~ ions.

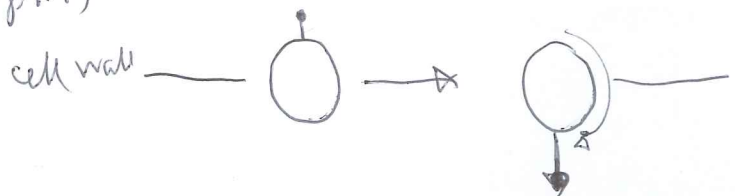
These exist both inside and outside the cell, creating a potential difference.

The cell walls are permeable - ions may be transported through the cell membrane, passing through pores called ~~from~~ channels or gates.

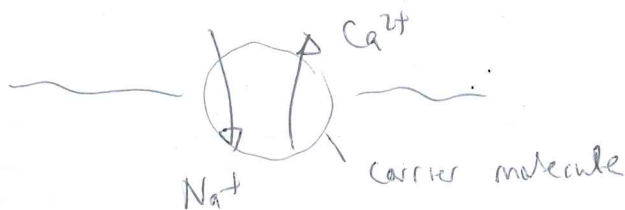
Osmosis is the mechanism by which ~~so~~ water is transported across the cell membrane.

Carrier mediated diffusion - a molecule hitches a lift by binding to a carrier molecule which is lipid soluble and can move through the membrane

Carrier mediated transport - a molecule binds to a protein that has an active site that may be exposed to the interior or exterior of the cell (eg glucose or amino acid transport)



Pumps - these exchange one ion for another eg Na^+ and K^+ or Na^+ and Ca^{2+}



A model for carrier mediated transport

C_i = state with binding site exposed to the exterior

C_e = state with binding site exposed to the interior

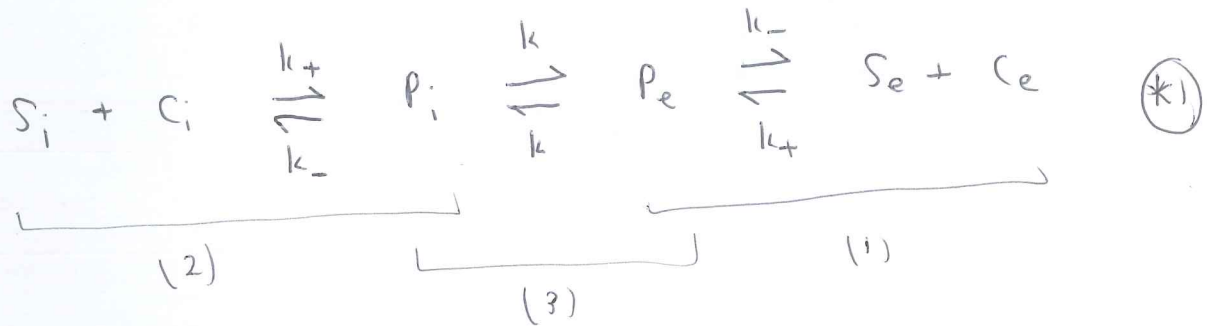
C_e can bind to make a complex P_e with a substrate molecule in the exterior, S_e , to make a product P_e (1)

C_i can bind to make a product P_i with a substrate molecule in the interior, S_i (with same rates as in the exterior) (2)

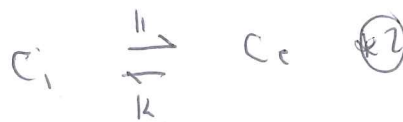
Further P_i can turn into P_e and vice versa (this is the carrier doing its 'rotation') (3)

See next page for pictures.

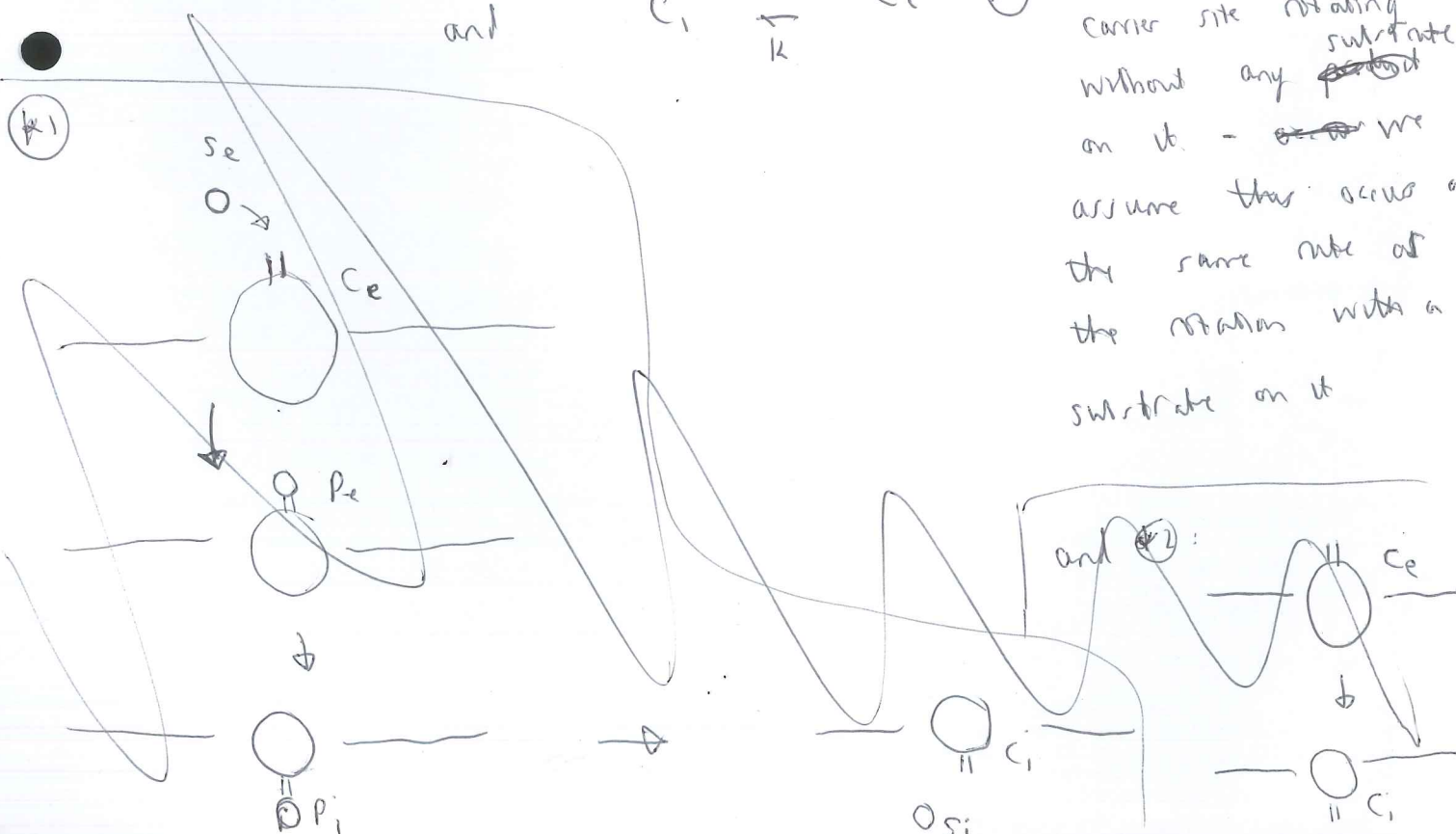
The reaction scheme is:



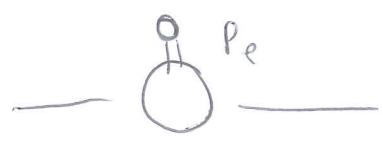
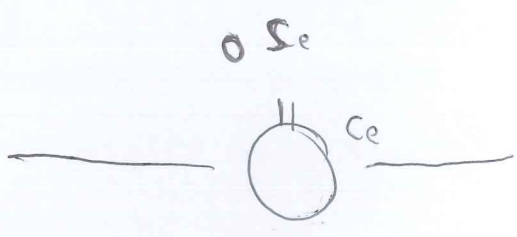
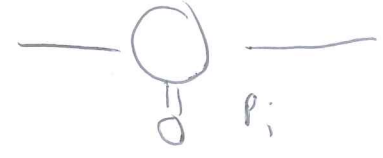
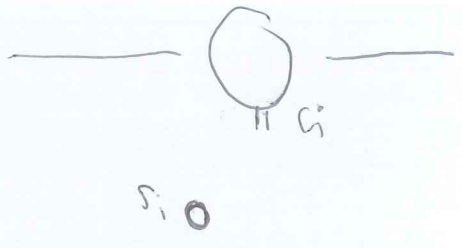
and



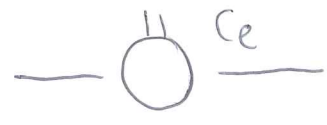
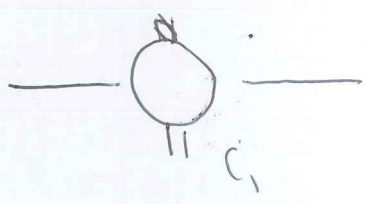
this is the carrier site rotating without any substrate on it - we assume this occurs at the same rate as the rotation with a substrate on it



(K1):



(K2):



Finally, assume that substrate is supplied from the exterior at a rate J_s and taken

Using the law of mass action we have

$$R_i = k_+ S_i C_i - k_- P_i$$

$$R_e = k_+ S_e C_e - k_- P_e$$

$$F = k P_i - k P_e$$

$$G = k C_i - k C_e$$

We are interested in finding a relationship for the rate of transfer of row from one side to the other in steady state and how this depends on the parameters in all the individual reactions

Finally, suppose that substrate is supplied from the exterior at a rate J and taken away from the interior at the same rate.

Then:

$$\frac{dS_i}{dt} = -J - R_i \quad (1)$$

$$\frac{dS_e}{dt} = J + R_e \quad (2)$$

$$\frac{dP_i}{dt} = R_i - F \quad (3)$$

$$\frac{dP_e}{dt} = F - R_e \quad (4)$$

$$\frac{dC_i}{dt} = -G - R_i \quad (5)$$

$$\frac{dC_e}{dt} = +G + R_e \quad (6)$$

If J is unknown then there is six equations for seven unknowns.

Adding all ~~these~~ ^{(3) + (4) + (5) + (6)} equations gives

$$\frac{d}{dt} (\cancel{S_i} + P_i + P_e + C_i + C_e) = 0$$

$$\Rightarrow \cancel{S_i} + P_i + P_e + C_i + C_e = \text{constant}$$

This is conservation of carrier.

~~If J is also unknown~~

And (1) + (2) + (3) + (4) gives $S_i + S_e + P_i + P_e = \text{constant}$

This is conservation of substrate.

One can solve ① - ① in steady state to find
(see problem sheet 1, question 3)

$$J = \frac{k_- k C_e}{2k_0} \frac{S_e - S_i}{(K_m + S_i)(K_m + S_e) - K_d^2}$$

$$K_m = \frac{k_- + k_+}{k_4}$$

$$K_d = \frac{k_c}{k_+}$$

so this tells us the flux of ions transported across the cell membrane in steady state

Note the similarity in structure to the Michaelis-Menten flux we derived.

$$\left(\frac{dJ}{dS} = \frac{-J(S)}{S(K)} \right)$$

Active transport - the sodium-potassium pump

The carrier mediated transport described above moves molecules down chemical gradients.

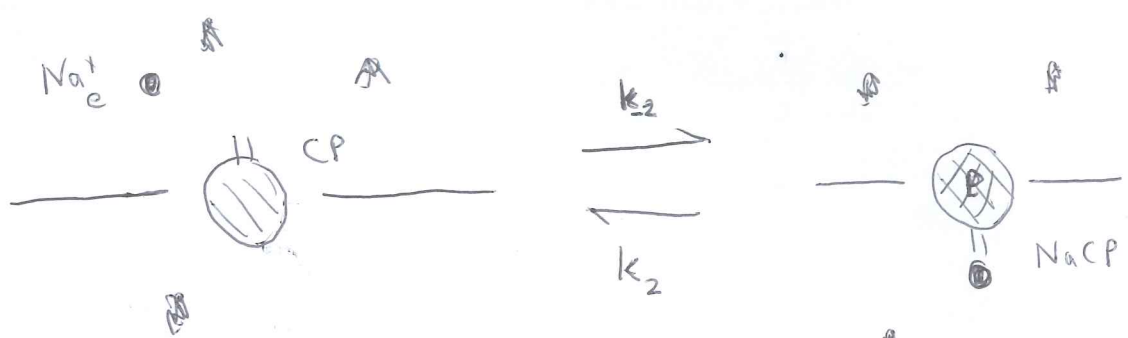
To move molecules against a chemical gradient requires energy. ~~and~~ This is known as an active transport mechanism.

One of the most important active transport mechanisms is the $\text{Na}^+ \text{K}^+$ pump.

For lectures just make the distinction from the carrier mediated transport before. Part on next page.



Done in this in lecture.

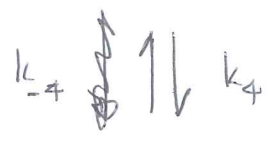
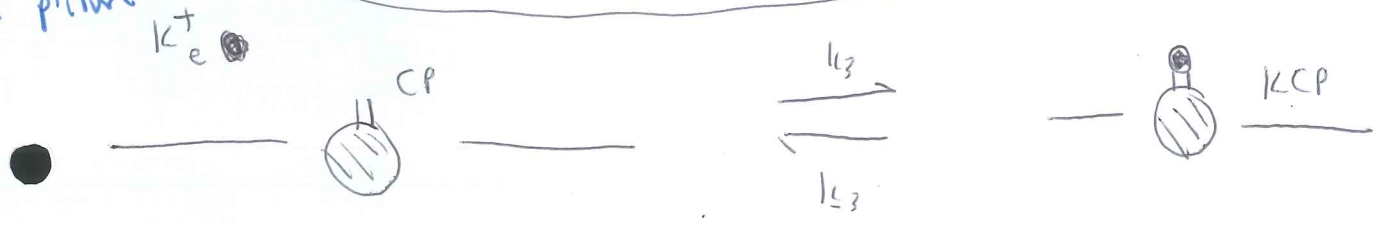


(requires energy) and provides phosphate

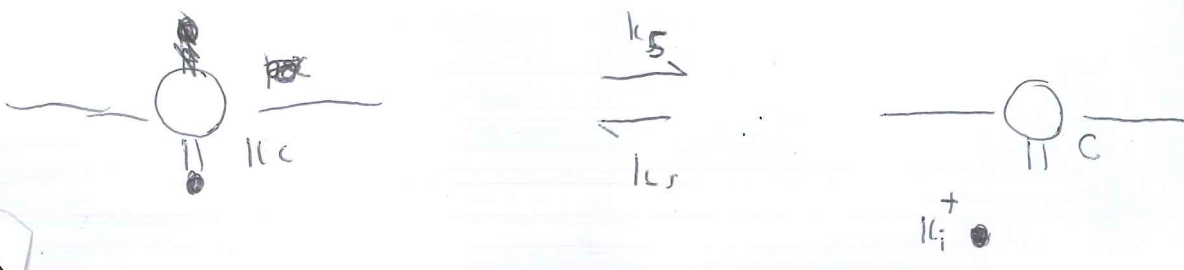
(in phosphorylated state which prefers to be pointing outward with a Na⁺ attached)

Could just say that and no need to delve into the picture

This is like the carrier mediated transport but now there is a chemical reaction which requires energy - this allows chemicals to move against concentration gradients



(phosphate ejected) and water

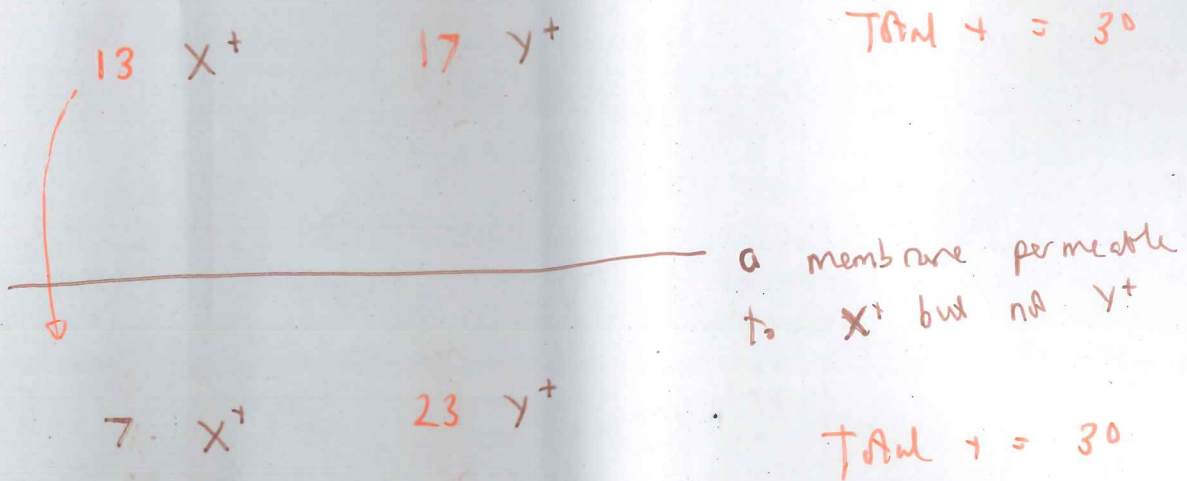


The Nernst potential and the resting potential

The Nernst potential is that obtained when all gates are open and there is a balance between diffusive flux and electric flux. The system is in equilibrium.

The resting potential is the difference between the potential outside and far from the cell and inside the cell, which may be different to the Nernst potential because gates open and close and ions are moved under different ionic and concentration gradients.

The Nernst potential

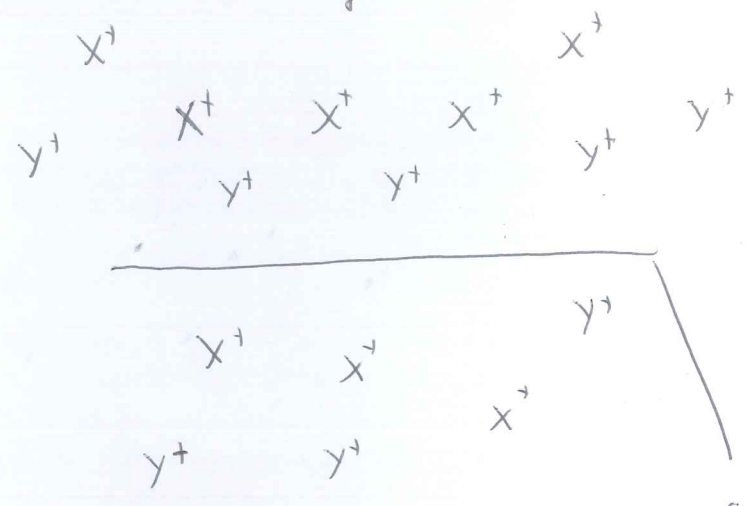


11 X^+ 17 Y^+

9 X^+ 23 Y^+

But doesn't reach 10 X^+ on each side because of electrostatic potential pushing back

The membrane potential
Consider the following set-up:



Equal positive and negative charges in each ~~side~~ respective side but different amounts on the two sides.

a membrane permeable to X^+ but not Y^+

X^+ will diffuse through the membrane to balance the charge of X^+ on both sides.

The balancing will happen in a small region near the membrane (so far from the membrane the ~~membrane~~ liquids will remain electro neutral.)

The potential difference that builds up ~~across~~ across the cell membrane as a result is called the Nernst potential

(ie. concentrations of X^+ will be equal across the membrane and concentrations)

What is this potential?

There will be a flux of ions due to a concentration gradient
There will be a flux of ions due to a potential difference

$J_1 = -D \nabla c$ (regular diffusive flux)

$d = \text{electric potential}$
 $\nabla d = \text{electric field}$

$J_2 = - \frac{u z c}{|z|} \nabla \phi$
 $u = \text{ion mobility} = \text{velocity under a constant electric field}$
 $z = \text{Valency of ion } (\frac{z}{|z|} = \pm 1 \text{ giving ion charge})$

So the total flux is $\underline{J} = \underline{J}_1 + \underline{J}_2$

$$\underline{J} = -D \nabla c - \frac{uzc}{|z|} \nabla \phi$$

or $J = -D \frac{\partial c}{\partial x} - \frac{uzc}{|z|} \frac{\partial \phi}{\partial x}$ (*) assuming set-up is one dimensional

Einstein's relation connects the diffusivity with the ion mobility:

$$D = \frac{uRT}{|z|F}$$

- R = universal gas constant
- T = absolute temperature
- F = Faraday's constant

(cf. Stokes Einstein:)

$$D = \frac{k_B T}{6\pi\eta a}$$

In equilibrium $J = 0$ (no flux)

We can integrate (*) to get

$$V = \phi_i - \phi_e = \frac{RT}{zF} \ln\left(\frac{c_e}{c_i}\right)$$

Nernst potential

e = exterior
i = interior

~~Now consider a~~

Ionic currents

The flow of ions across a membrane causes a build up of charge which means the membrane acts as a capacitor.

The voltage across the membrane is $V = \frac{Q}{C}$ (charge difference / capacitance)

The current across the membrane is $I = -\frac{dQ}{dt}$

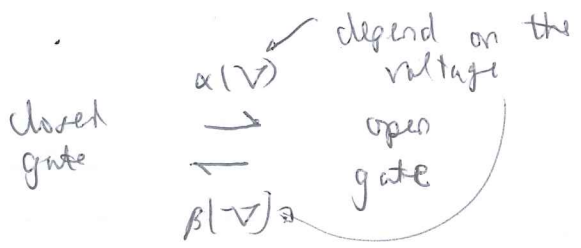
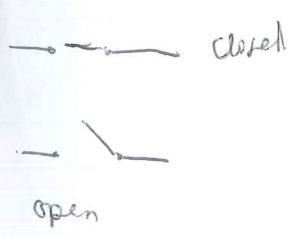
Combining these gives $C \frac{dV}{dt} + I = 0$ (assuming C = constant)

This equation is the basis for much theoretical electrophysiology

We link the current I_s to the voltage via $I_s = g_s(V - V_s)$ (ion-specific membrane conductance)

The total current $I = \sum I_s$ (just $V = IR$ and conductance is $\frac{1}{R}$)

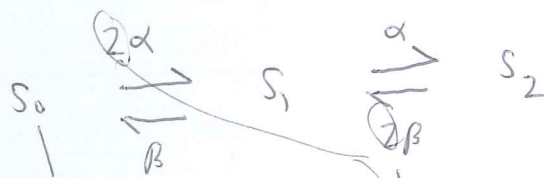
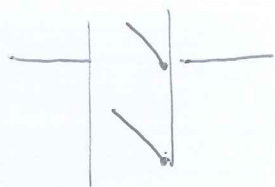
Gates But what does g_s look like? This is nA a constant - it depends on the fraction of gates that are available that are open, n. Then $g_s = n g_{s,max}$



conductivity when all gates are open

Law of mass action gives $\frac{dn}{dt} = \alpha(V)(1-n) - \beta(V)n$ (where $n_{\infty} = \frac{\alpha(V)}{\alpha(V) + \beta(V)}$ and $\tau = \frac{1}{\alpha(V) + \beta(V)}$)
 or $\frac{dn}{dt} = n_{\infty} - n$
 function of open gates

You can also have multiple gates:



number of closed gates

factor of 2 because there are two configurations for going from 2 open to 1 open gate and similarly for closed to open

Law of mass action gives $\frac{dS_0}{dt} = \beta S_1 - 2\alpha S_0$

$$\frac{dS_1}{dt} = \alpha S_0 - 2\beta S_1$$

and $\frac{dS_2}{dt}$ equation or just infer $S_0 + S_1 + S_2 = 1$ (conservation of gates)

We can reduce to one ODE by ~~finding~~ finding solution is $S_0 = (1-n)^2$, $S_1 = 2n(1-n)$, $S_2 = n^2$

where n satisfies $\frac{dn}{dt} = \alpha(1-n) - \beta n$ (i.e. the one-gate equation)

[Problem Sheet 2, question 1]

In this case the proportion of open channels is $S_2 = n^2$ so the conductivity in this case is $n^2 g_{max}$.

This result generalizes for N gates with a conductivity of $n^N g_{max}$

You can also look at the case of non-identical gates

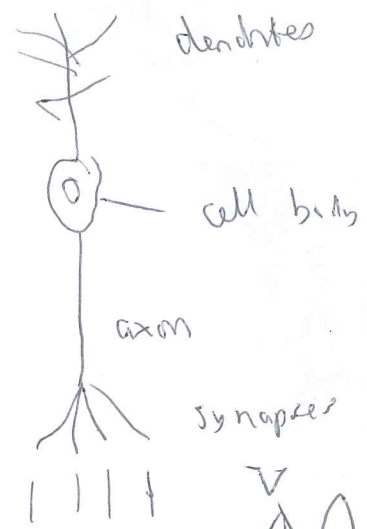
You can also look at the case of non-identical gates - see lecture notes page 39 and problem sheet 2 question 8.1.

The Hodgkin-Huxley model

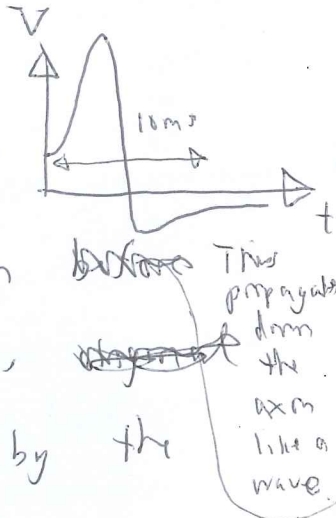
The nervous system is a communication system formed by nerve cells or neurons. Information is propagated along long cylindrical segments called axons by electrochemical signals.

Communication between cells occurs at junctions between synapses ~~and~~ dendrites.

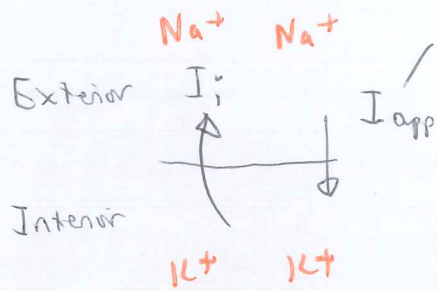
If a small current is applied for a short time then the membrane potential just returns to its resting potential when the current is removed.



But for a sufficiently high current, the membrane potential undergoes a large excursion ~~before~~ ~~returning~~ before returning to its resting value. Signals are transmitted by the propagation of these action potentials down an axon. Later on we will look at this spatial propagation but for now we



will look at a spatially independent model. ~~This~~ This can be achieved in practice by inserting an electrode along the axon to spread the current out - this is called the space clamp technique.



Apply a current I_{app} to the axon and observe the ^{ionic} current that comes out, I_i .

Our earlier equation

$$C \frac{dV}{dt} + I = 0 \quad (1)$$

$$\Rightarrow C_m \frac{dV}{dt} + (I_i - I_{app}) = 0.$$

Capacitance of the ~~axon per unit area~~ membrane

Potassium flow

Recall we have

our link between V and I_i :



$$I_i = g_s (V - V_s)$$

and $g_s = \cancel{g_s} N g_{smax}$

where N = number of gates

and $\tau(V) \frac{dn}{dt} = n_{\infty}(V) - n.$ (2)

$$g_s = N^4 \frac{g_{smax}}{2N_0}$$

What we find is that the potassium conductance may be controlled by this model when $N=4$. Note, however that this is not due to four gates but just an experimental fit.

n is called the potassium activation

For the sodium conductance a model of the form turns the sodium current on and another which turns it off. This can be described by

$$I_s = g_{Na} m^3 h$$

$$\tau_m(V) \frac{dm}{dt} = m_{\infty}(V) - m \quad (3)$$

$$\tau_h(V) \frac{dh}{dt} = h_{\infty}(V) - h \quad (4)$$

Again, these time constants are experimental fit rather than related to gates

is appropriate. This is like two gates

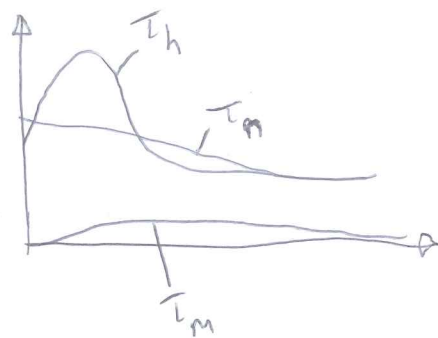
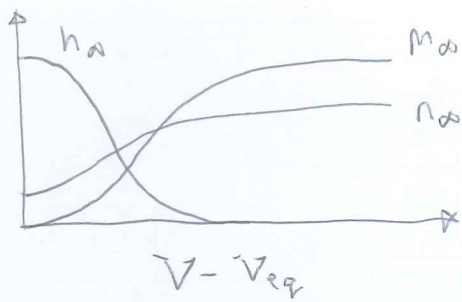
m is called the sodium activation
 h is called the sodium inactivation.

The Hodgkin-Huxley model for the current is then

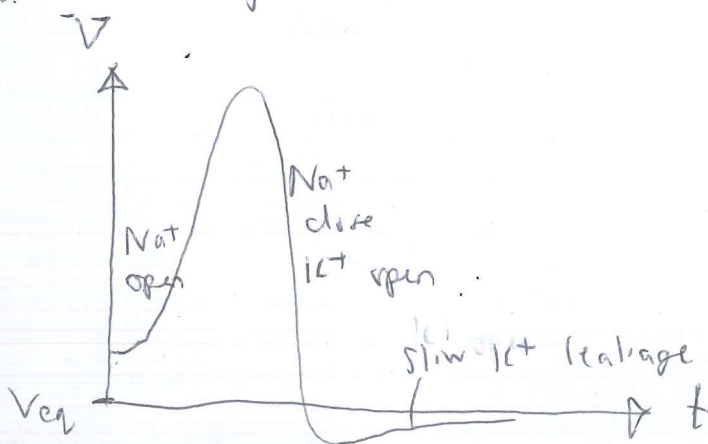
$$I_i = \underbrace{g_{Na} m^3 h (V - V_{Na})}_{Na^+ \text{ current}} + \underbrace{g_K n^4 (V - V_K)}_{K^+ \text{ current}} + \underbrace{g_L (V - V_L)}_{\text{leakage (mainly Cl- chloride ions)}}$$

(This model also comprises ~~equations~~ the DPE system (1)-(4))

What does the potential do?



- 1) Apply a current I_{app} .
- 2) V rises due to the current (equation 1)
- 3) τ_m is small so m rises quickly (equation 3)
 $\Rightarrow Na^+$ floods into the axon from outside. (since $g_r = m^3 h g_{Na}$)
- 4) This causes even more of a potential difference, causing m_{∞} to rise (graph 1), which ~~this~~ causes even more of a potential difference.
- 5) m_{∞} falls causing Na^+ flow to slow as g_{Na} falls
 n_{∞} rises causing K^+ to flow out of (gate closer)
 the axon (g_{K^+} rises - gate opens) $g_s = n g_{K^+}$
 This causes an overshoot past V_{eq}
- 6) Slow leakage of K^+ back into the axon causes the system to go back to equilibrium.



The membrane is excitable

The equilibrium is a steady state but a large enough perturbation (which in practice isn't too large) causes the potential to undergo an excursion - the action potential.

At the moment this is a low dimensional system (V, n, m, h described by equations ①-④). This is ^{not} easy to analyse but we can reduce this to a two-dimensional system called the Fitzhugh-Nagumo

~~equations~~ model

The FitzHugh Nagumo model

Assumptions: 1) τ_m is small so $m \approx m_\infty(V)$
 (m rapidly reaches its quasi-stable value, using ③)

2) $\tau_n = \tau_h$ (not perfect, see graph 2, but a decent approximation)

3) $n_\infty + h_\infty = \text{constant}, \bar{h}$ (motivated by graph 1)
 $\Rightarrow n + h = \bar{h}$ (using ② and ④)

This reduces the Hodgkin-Huxley model (four ODEs for V, n, m, h) to the two-dimensional system for V and n :

$$C_m \frac{dV}{dt} = I_{\text{app}} - (g_K(V - V_K)n^4 + g_{Na}(V - V_{Na})m_\infty^3(V)(\bar{h} - n) + g_L(V - V_L))$$

$$\tau_n(V) \frac{dn}{dt} = n_\infty(V) - n$$

see problem sheet 2 for this.

$$\frac{d}{dt}(n+h) = \bar{h} - (n+h)$$

$$\Rightarrow (n+h) = \bar{h} + ce^{-t}$$

This system is true for all time t eventually this settles to $n+h = \bar{h}$.

Non-dimensionalization

$$v = \frac{V - V_{eq}}{V_{Na} - V_{eq}}$$

$$t = \tau_n t'$$

resting potential for Na⁺

$$\begin{aligned} I^+ &= \frac{I_{app}}{g_{Na} V_{Na}} & V_{K}^+ &= \frac{-V_K}{V_{Na}} \\ \gamma_K &= \frac{g_K}{g_{Na}} & V_L^+ &= \frac{V_L}{V_{Na}} \\ \gamma_L &= \frac{g_L}{g_{Na}} & \epsilon &= \frac{C_m}{g_{Na} \tau_n} \end{aligned}$$

$$\Rightarrow \begin{cases} \frac{dn}{dt} = n_{\infty}(v) - n & \textcircled{1} \\ \epsilon \frac{dv}{dt} = I^* - g(v, n) & \textcircled{2} \end{cases}$$

$$g(v, n) = \gamma_K (v + v_K^+) n^4 + \gamma_L (v - v_L^+) - (1-v)(h-n)m^3(v)$$

This is why you non-dimensionalize - to see the relative parameter sizes

key point: $\epsilon \ll 1$

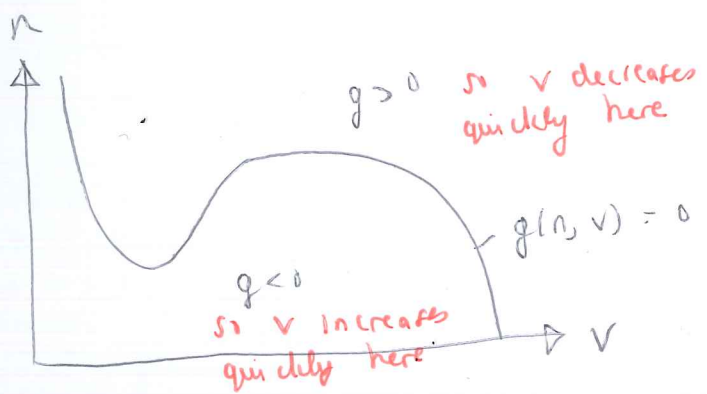
quicker reaches quasistatic equilibrium, in $\textcircled{2}$ and $\gamma_L \ll 1$, so $g(v, n)$ simplifies

Phase plane analysis

start by considering the case $I^* = 0$.

Setting $\epsilon = \gamma_L = 0$ in $\textcircled{2}$ gives $\frac{n^+}{h-n} = \frac{(1-v)m^3(v)}{\gamma_K(v+v_K^+)}$

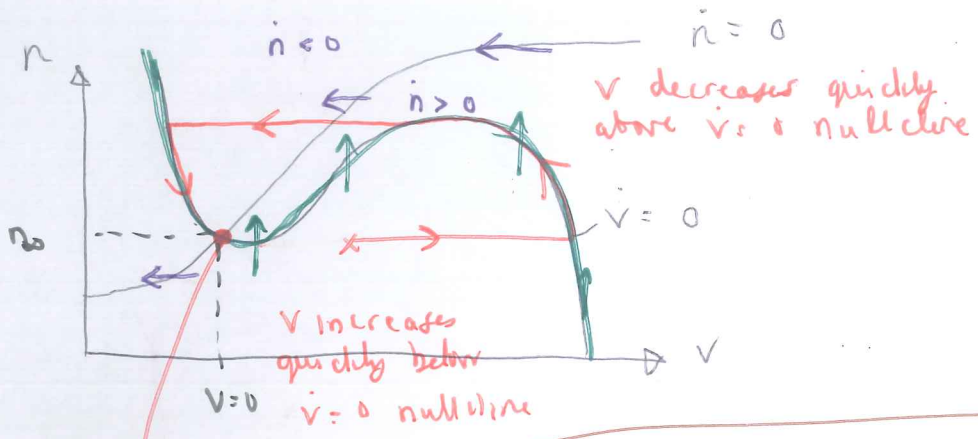
an algebraic relationship between n and v



Since $\epsilon \ll 1$ we know that the system quickly jumps onto this manifold

Note that including the γ_L term doesn't change these qualitative features

Now we just need to add the $\dot{n} = 0$ nullcline
 ($n = n_0(v)$)



Proved below so don't need to say this

Reaches this (unique) fixed point.

Perturbing around the fixed point we see that trajectories spiral around so the fixed point is a spiral.

Although this fixed point is stable, only a small increase in v will lead to a large excursion - this is the action potential again. If we plotted v versus t we would obtain the graph we drew earlier.

Stability

Linearize near the fixed point

Set $v = v_0 + \delta v$
 $n = n_0 + \delta n$

and ODEs become

$$\frac{d}{dt} \begin{pmatrix} \delta n \\ \delta v \end{pmatrix} = \begin{pmatrix} -1 & \frac{dn_0}{dv} \\ -\frac{dg}{dv} & -\frac{dg}{dv} \end{pmatrix} \begin{pmatrix} \delta n \\ \delta v \end{pmatrix}$$

Fixed point is $n = 0, v = 0$

$\Rightarrow n = n_0, v = 0$

(By definition since v is the potential relative to the equilibrium)

Stability is given by $\det(M)$ and $\text{tr}(M)$

$\det(M) < 0 \Rightarrow$ saddle

$\det(M) > 0$ and $\text{tr}(M) < 0 \Rightarrow$ node or spiral

(See next page for why.) trace

Trace and determinant method

Eigenvalues can be written as

$$\lambda = \frac{\text{Tr}(\underline{M}) \pm \sqrt{(\text{Tr}(\underline{M}))^2 - 4 \det(\underline{M})}}{2}$$

We have $\det(\underline{M}) = \frac{1}{3} \left(\left. \frac{\partial g}{\partial n} \right|_{n=n^*} + \left. \frac{\partial g}{\partial v} \right|_{n=n^*} \frac{dn_{ss}}{dv} \right)$

New proof
 View $\det(\underline{M}) > 0$
 slope of n nullcline = $\frac{dn_{ss}}{dv}$

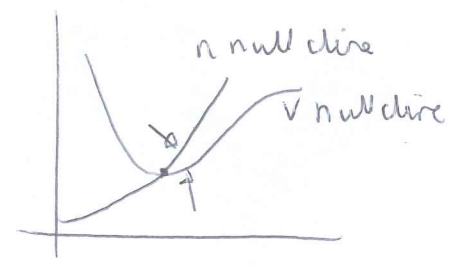
slope of v nullcline = $-\frac{\frac{\partial g}{\partial n}}{\frac{\partial g}{\partial v}}$

since $g(v, n) = 0$

$$\frac{\partial g}{\partial n} \frac{dn}{dv} + \frac{\partial g}{\partial v} = 0$$

$$\frac{dn}{dv} = - \frac{\frac{\partial g}{\partial n}}{\frac{\partial g}{\partial v}}$$

from graph, slope of n nullcline > slope of v nullcline



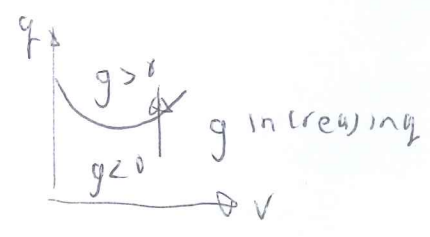
$$\frac{dn_{ss}}{dv} > - \frac{\frac{\partial g}{\partial n}}{\frac{\partial g}{\partial v}}$$

$$\det(\underline{M}) > 0$$

$$\text{tr}(\underline{M}) = -1 - \frac{1}{3} \left. \frac{\partial g}{\partial v} \right|_{n=n^*}$$

Fixed point is stable if $\text{tr}(\underline{M}) < 0 \Rightarrow \frac{\partial g}{\partial v} > -3$

$\frac{\partial g}{\partial v} > 0$ because of picture



fixed point is stable

27
Although the fixed point is stable, a small increase
in v will lead to a large excursion - this is
the action potential again. If we plotted v versus
 t we would obtain the graph we drew earlier.

Limit cycles

If we apply a current then this will push us off the equilibrium point and send us round the trajectory before starting the process all over again - we only need a bit of energy to achieve this.

Slightly different to a conventional limit cycle because in this case you need

The FitzHugh-Nagumo model is the reduction of the four dimensional Hodgkin-Huxley model to a two-dimensional system.

a bit of energy to kick it round the cycle

The FitzHugh-Nagumo equations are an analytically similar pair of equations that have the same behaviour.

$$\epsilon \dot{v} = J^+ + f(v) - w$$

$$\dot{w} = \gamma v - w$$

$$\bullet \quad f(v) = v(v-a)(1-v) \quad 0 \leq a \leq 1$$

FitzHugh-Nagumo equations

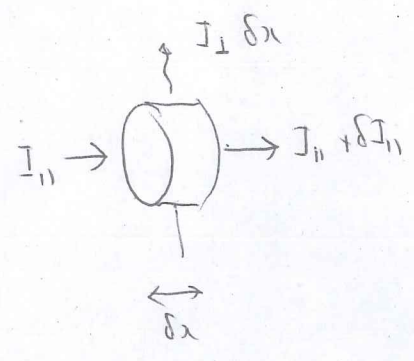
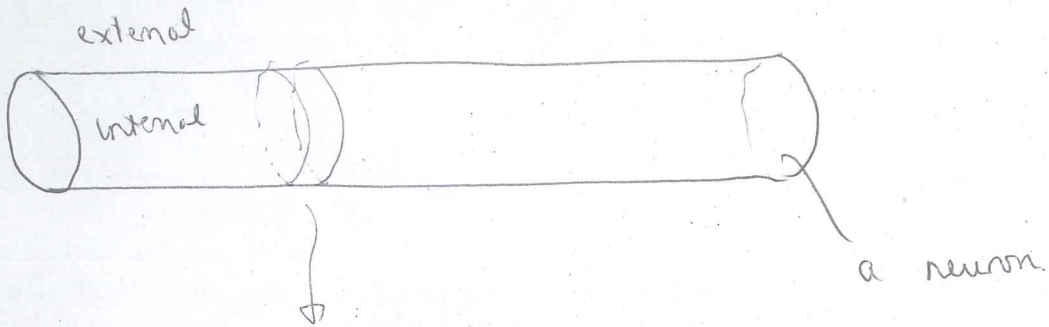
$$\epsilon \dot{v} = J^* - g(v, n)$$

$$\dot{n} = n_\infty(v) - n$$

FitzHugh-Nagumo model

Wave propagation in neurons

We now explore spatial dependence of the Hodgkin-Huxley model



- I_{\perp} = transmembrane current per unit length
- $I_{||}$ = axial current
- R = axial resistance
- C = capacitance per unit length

In a segment δx the total charge is $C V \delta x$

Charge conservation:

$$\frac{\partial}{\partial t} (C V \delta x) = - I_{\perp} \delta x + I_{||} - (I_{||} + \delta I_{||})$$

current left through wall
current in left
current out right

$$C \frac{\partial V}{\partial t} \delta x = - I_{\perp} \delta x - \frac{\delta I_{||}}{\delta x} \delta x$$

assuming $C = \text{constant}$

$$C \frac{\partial V}{\partial t} = - I_{\perp} - \frac{\partial I_{||}}{\partial x} \quad (*)$$

$$-\delta V = I_{||} R \delta x$$

Definition of resistance
(ie $\Delta V = I_{||} R_{total}$)

$$\Rightarrow \frac{\delta V}{\delta x} = -I_{||} R$$

So in (*),

$$\boxed{C \frac{\partial V}{\partial t} = -I_{\perp} + \frac{1}{R} \frac{\partial^2 V}{\partial x^2}} \quad (7)$$

This is called the telegraph equation
or the cable equation.

If the ~~rise~~ ^{neuron} perimeter is $p = \pi d$
diameter

$$\text{then } I_{\perp} = p(I_i - I_{app})$$

current per unit area between inside and outside as defined earlier

and $C = p C_m$
|
actual capac
capacitance per unit area.

If R_c = resistivity of medium then $R = \frac{R_c}{A}$

where $A = \frac{1}{4} \pi d^2$ is the neuron cross-sectional area.

In (7) this gives
$$C_m \frac{\partial V}{\partial t} = I_{app} - I_i + \frac{d}{4R_c} \frac{\partial^2 V}{\partial x^2}$$

Non-dimensionalization

$$v = \frac{V - V_{eq}}{V_{Na} - V_{eq}}, \quad I_i = g_{Na} (V_{Na} - V_{eq}) \cdot g(l_n, v), \quad x = l \hat{x}$$
$$t = \tau_n \hat{t}$$

where l is to be chosen later.

(Same as earlier non-dimensionalization)

This gives

$$\epsilon \frac{\partial v}{\partial t} = I^* - g(l_n, v) + \epsilon^2 \frac{\partial^2 v}{\partial x^2}$$

$$\frac{\partial n}{\partial t} = n_\infty(v) - n$$

(see problem sheet 2 for derivation of this)

This is the space dependent version of the ~~Hodgkin-Huxley~~ ^{FitzHugh-Nagumo} model - it's the same but with a $\frac{\partial^2 v}{\partial x^2}$ term.

Let's analyse action potentials in this case. We could analyse the equations above but it is easier to analyse the space-dependent FitzHugh-Nagumo equations (the equations that display the same qualitative behaviour but are easier to analyse):

$$\epsilon \frac{\partial v}{\partial t} = f(v) - w + \epsilon^2 \frac{\partial^2 v}{\partial x^2}$$

$$\frac{\partial w}{\partial t} = \gamma v - w$$

$$f(v) = v(v-a)(1-v)$$

and γ large enough that $(0,0)$ is the unique steady state.

Let's look for a travelling wave solution so we can find a wave travelling down the axon:

$v = v(\xi), w = w(\xi), \xi = ct - x, c > 0.$

$\epsilon c v' = f(v) - w + \epsilon^2 v''$

$cw' = \gamma v - w.$

with $v, w \rightarrow 0$ as $\xi \rightarrow \pm\infty.$

It is harder to do phase plane analysis now because the phase plane is three dimensional rather than two:

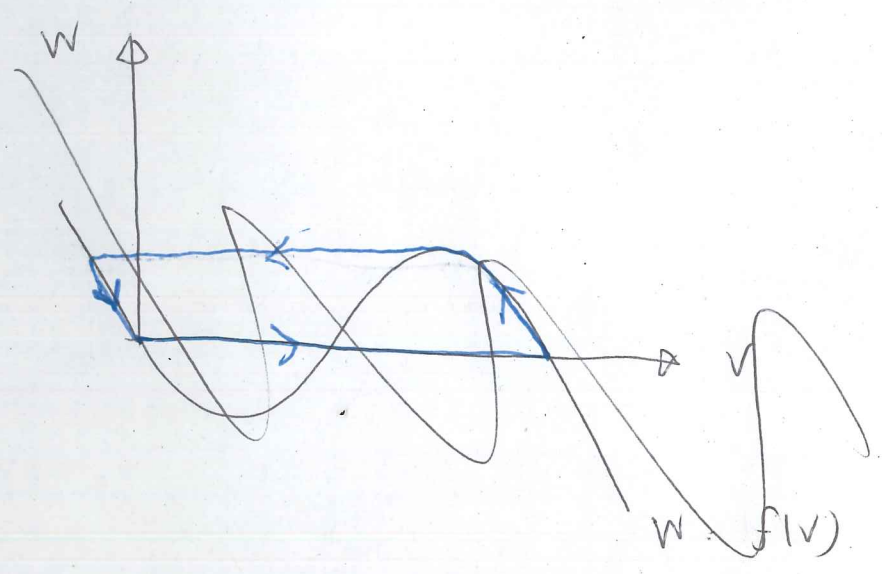
v, w, v'

However, $\epsilon \ll 1$ so this allows us to make progress without having to consider the three-dimensional phase space.

~~If we ignore the ϵ^2 then we have a picture like~~

~~below:~~

There are four different regions of behaviour.



To begin with, if we are not on the curve $w = f(v)$ then we quickly move there because of

$$\epsilon v' = \underbrace{f(v) - w}_{\neq 0} + \epsilon^2 v''$$

fast motion
higher order correction
just like before (ignoring higher order ϵ^2 term)

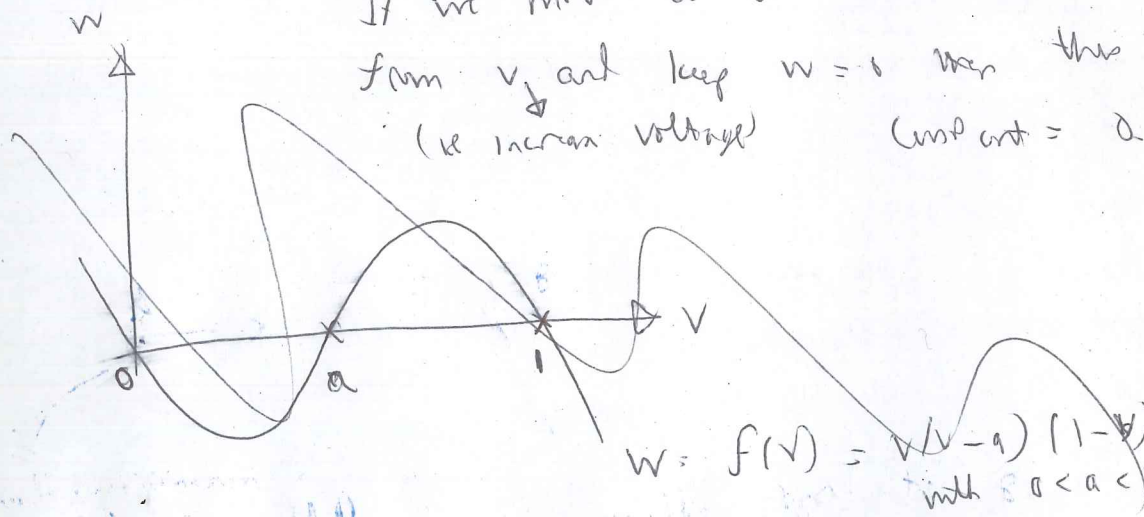
In this region, things happen over a fast ξ scale. This suggests rescaling $\xi = \epsilon \zeta$

$$\epsilon c \frac{dv}{d\zeta} = f(v) - w + \frac{d^2 v}{d\zeta^2}$$

and in this region, considering the other equation,

$$w' = \gamma v - w \Rightarrow c \frac{dw}{d\zeta} = \epsilon (\gamma v - w) \Rightarrow w = \text{constant}$$

If we move a little bit away from v_1 and keep $w = 0$ then this (is increase voltage) constant = 0



$$w = f(v) = v(v-a)(1-v)$$

with $0 < a < 1$

Fixed points of this rescaled system are $v = 0, a, 1$. Stability analysis (is linearization) shows that $v = 0, 1$ are saddles and $v = a$ is an unstable node or spiral (see lecture notes)

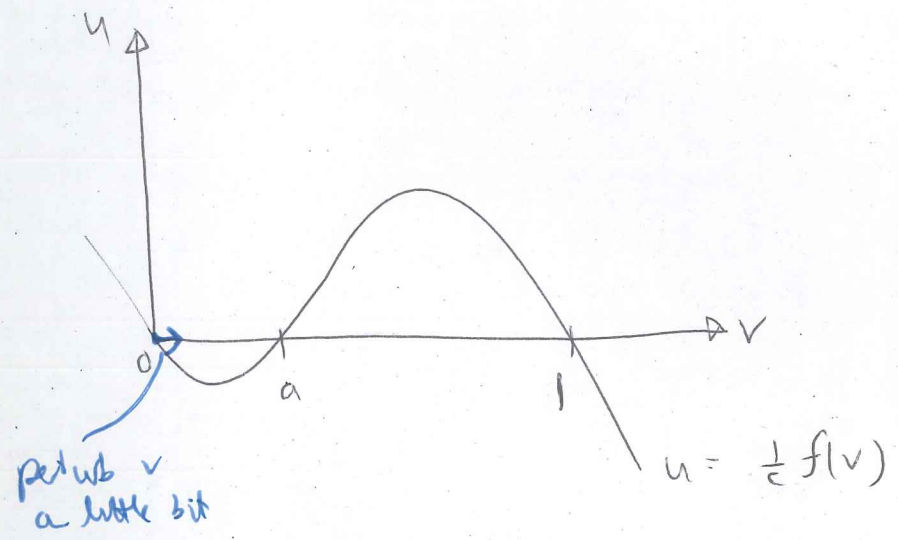
Now set $\frac{dv}{dt} = u$

Then our phase plane is $u' = u$

$u'' = cu - f(v)$
" $v(v-a)(1-v)$
 $0 < a < 1$

~~Now only one trajectory to~~

Fixed points of this reduced system are $u=0, v=0, a, 1$

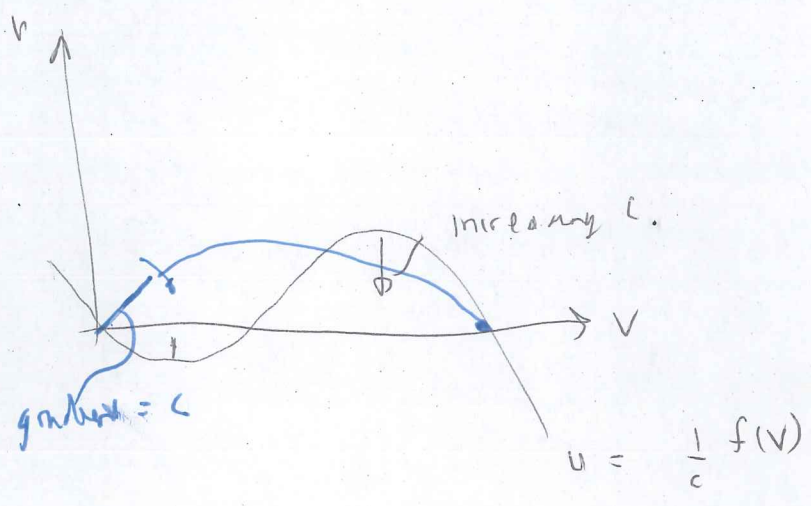


Linear stability analysis shows that $v=0, 1$ are saddles and $v=a$ is an unstable node. So we are interested in the trajectory in the phase plane that goes from $v=0$ to $v=1$ fixed points (to ~~replicate~~ replicate the action potential we had in the space-clamped case where we had the fast behaviour jumping out the nucleus)

There is only one value of c that achieve this.

Now $\frac{dn}{dv} = \frac{d \frac{u'}{v}}{v} = \frac{c u - f(v)}{u} = c$ at $v=0$ so gradient of

trajectory is c



Like a shooting problem:

This is how c is selected - this means there is a unique wave speed for the travelling wave.

(from $n' = c u - f(v)$)

(ii) Then, once we land on the $n=0$ nullcline $u = \frac{f(v)}{c}$ we slowly move up this. On this,

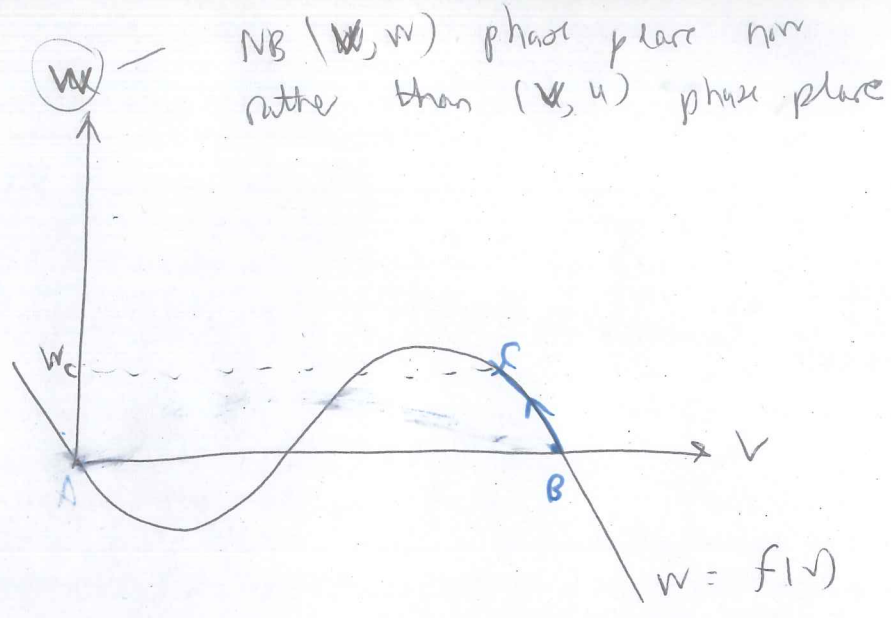
$$\epsilon c v' = f(v) - w + \epsilon^2 v''$$

$$c w' = \gamma v - w \quad (2)$$

$$\epsilon f(v) = f(v) - w + \epsilon^2 v''$$

$$\Rightarrow w \approx f(v) \quad (1)$$

This takes us up the curve $u = \frac{f(v)}{c}$ until we reach $w = \gamma v$ (the equilibrium of (2))



Note that ζ is no the maximum of $w = f(v)$ unlike in the space-damped model. Now ζ is where $w = \gamma v$. We need to find what this value (w_c) is, which we will find out in the next stage.

(iii). Once we have reached this point we enter another fast phase. Again rescale $\xi = \epsilon \zeta$ to capture this

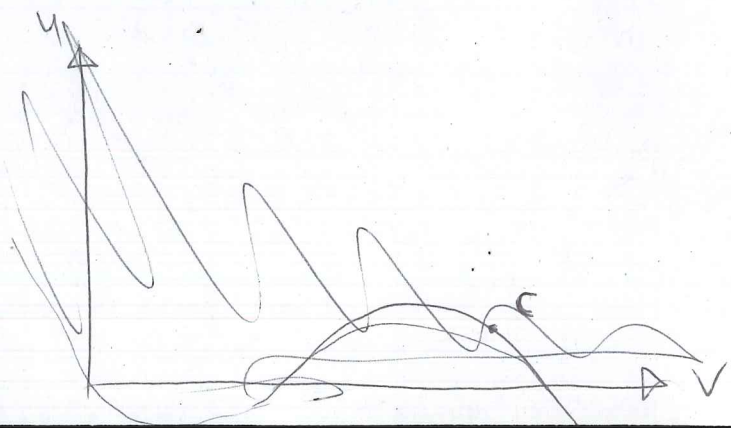
but this time $w = w_c$ (a non zero constant). Then the

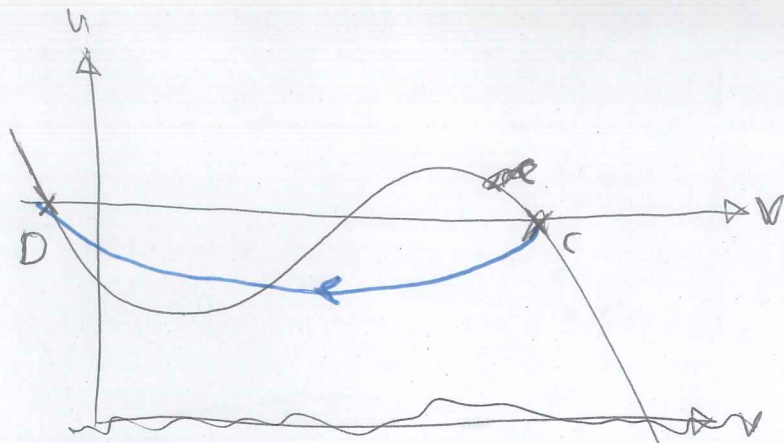
system is

$$c v' = f(v) - w_c + v''$$

which we need to find out what value it is

$$\begin{aligned} \dot{v} &= u \\ u' &= c u - f(v) + w_c \end{aligned}$$



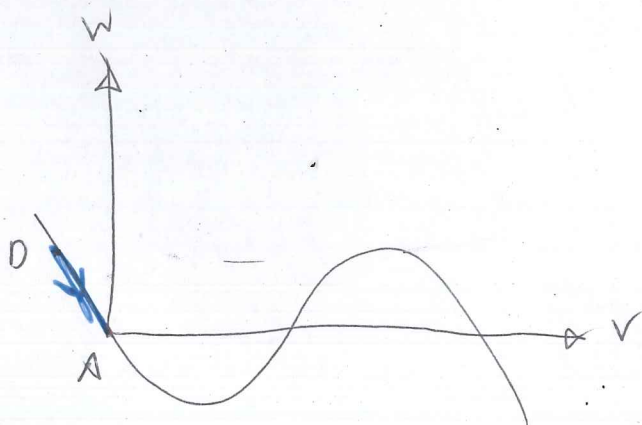


$$u = \frac{f(v)}{c} - \frac{w}{c}$$

(same curve as before just shifted down)

C and D are saddles and this time we have a trajectory that takes us from C to D. This time, to make sure these trajectories join up, it is we that we need to choose correctly (just like we had to choose c correctly in phase (i))

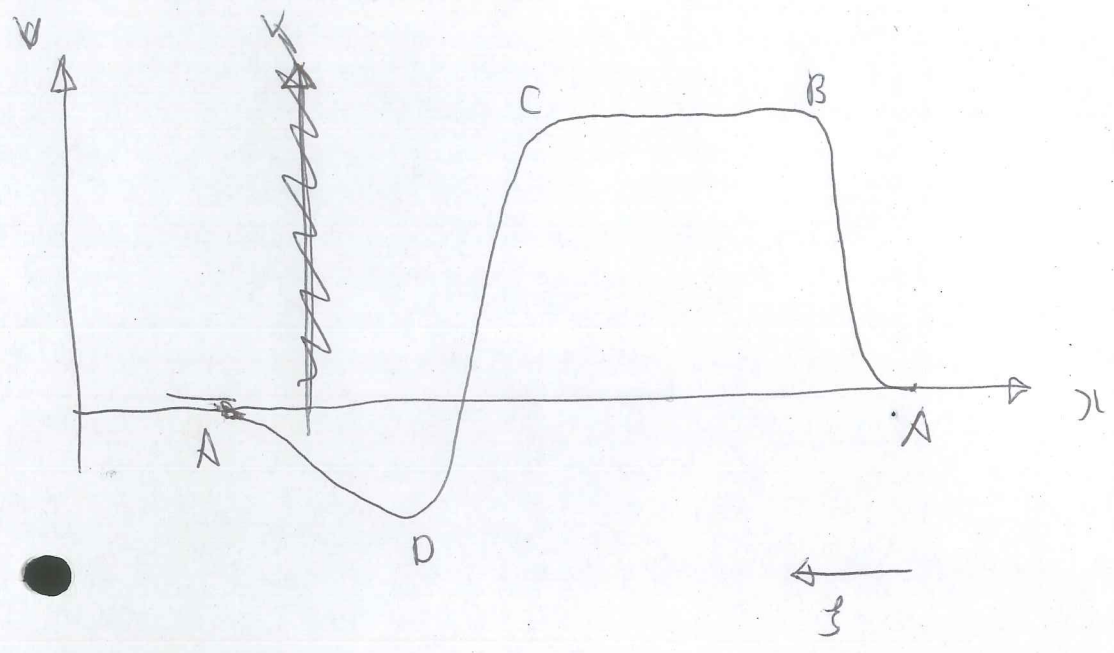
Finally, a slow phase takes us back to A again, on the (v, w) phase plane:



Again, on here,
 $w = f(v)$

$$w' = \gamma v - w \quad (< 0 \text{ so motion is downwards, back to A})$$

The overall picture is a travelling wave that moves down the axis and looks like this:



Reel from right to left

Calcium dynamics

Calcium (Ca^{2+}) is important in muscle dynamics and cell signalling.

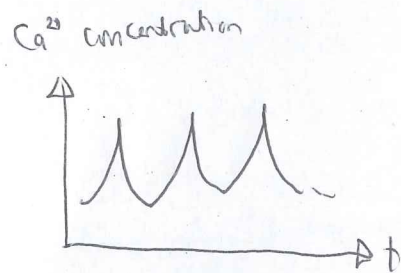
Ca^{2+} is stored in a ^{cell in} bones and released by hormonal stimulation. The internal store is called the sarcoplasmic reticulum. It releases Ca^{2+} via calcium induced calcium release.

Extracellular Ca^{2+} concentrations are higher than intracellular concentrations so Ca^{2+} must be pumped out. The intracellular fluid matrix is called the sarcoplasm.

Muscle cells are bundles (fascicles) of muscle fibres (cells) each of which contains arrays of filament structures (microfibrils) which contract under the action of Ca^{2+} .

Under stimulation from a nerve cell, an action potential is triggered and propagates along the fibre as we have seen last lecture. Na^+ floods in and this allows Ca^{2+} in too.

The release of Ca^{2+} is quite spiky:



Can we devise a mathematical model for muscle contraction with a low Ca^{2+} concentration that is excitable under stimulus? in steady state

The two-pool model

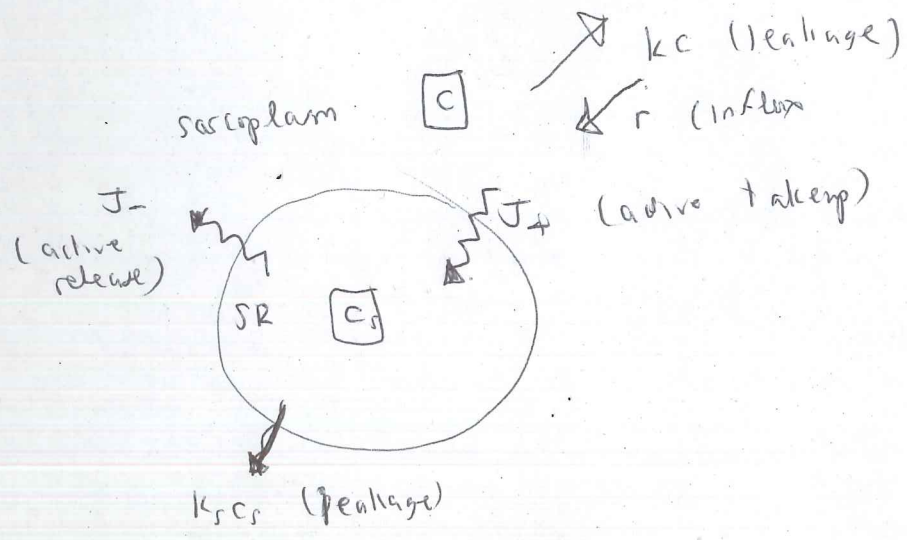
We want to derive a model ~~set~~ to explain how Ca^{2+} moves between the sarcoplasmic reticulum (the store) and the sarcoplasm.

- c = concentration of Ca^{2+} in the sarcoplasm
- c_s = concentration of Ca^{2+} in the sarcoplasmic reticulum (SR)
- J_+ = rate of ~~release~~ take-up of Ca^{2+} from by the sarcoplasmic reticulum (SR) (by receptors) [active uptake]
- J_- = rate at which the SR releases its internal store (calcium-induced calcium release) [active release]
- r = inflow of Ca^{2+} into the sarcoplasm, because of an applied stimulus (from the outside world)

~~k_{cs} = rate of leakage of Ca^{2+} from sarcoplasm~~ (passive - proportional to concentration)

k_{sr} = rate of leakage of Ca^{2+} from SR (passive - proportional to concentration) into the sarcoplasm

k_c = rate of leakage of Ca^{2+} from sarcoplasm to outside world (no path into the SR) (passive - proportional to concentration)



$$\frac{dc_s}{dt} = J_+ - J_- - k_c c_s \stackrel{\text{def}}{=} F$$

$$\frac{dc}{dt} = r - k_c c - (J_+ - J_- - k_c c_s)$$

$$= r - k_c c - F$$

We choose $J_+ = \frac{V_1 c^n}{K_1^n + c^n}$

NB V_1 not a voltage - this is a concentration rate

V_1, K_1, n our numbers

This is a Hill function

These bits are important

$J_- = \frac{V_2 c_s^m}{K_2^m + c_s^m} \left(\frac{c_f}{K_2^m + c_f} \right)$

This is the important bit that causes the calcium induced calcium release

Non-dimensionalization

$$c = K_1 u, \quad c_s = K_2 v, \quad t = \frac{1}{k} \hat{t}, \quad F = V_2 f$$

$$\frac{du}{dt} = N - u - \frac{\gamma}{\epsilon} f(u, v)$$

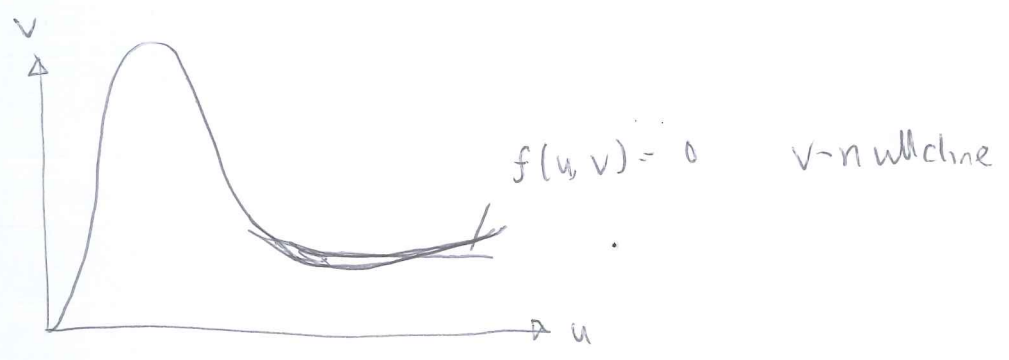
$$\frac{dv}{dt} = \frac{1}{\epsilon} f(u, v)$$

$$f = B \left(\frac{u^n}{1 + u^n} \right) - \left(\frac{v^m}{1 + v^m} \right) \left(\frac{u^p}{\alpha + u^p} \right) - \delta v$$

$$N = \frac{r}{k K_1}, \quad \gamma = \frac{K_2}{K_1}, \quad \epsilon = \frac{k_1 K_2 \ll 1}{V_1}, \quad \alpha = \frac{K_2}{K_1}, \quad B = \frac{V_1}{V_2}, \quad \delta = \frac{k_5 K_2 \ll 1}{V_2}$$

This is a two-dimensional system (u, v) so we may do phase-plane analysis

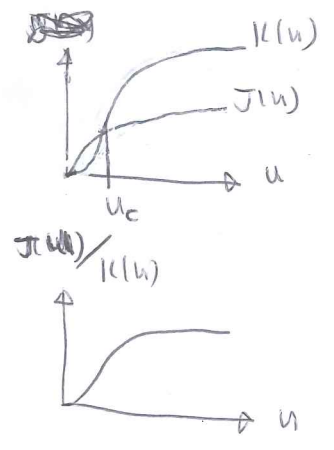
$\epsilon \ll 1$ means that we quickly jump onto the v -nullcline, $f(u, v) = 0$



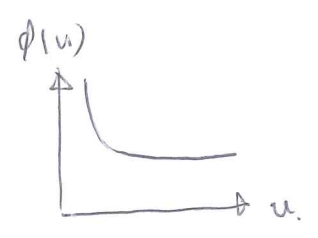
to plot this curve

1) $\delta \ll 1$ so ignoring the δ term in ~~the equation~~ $f(u, v)$ gives

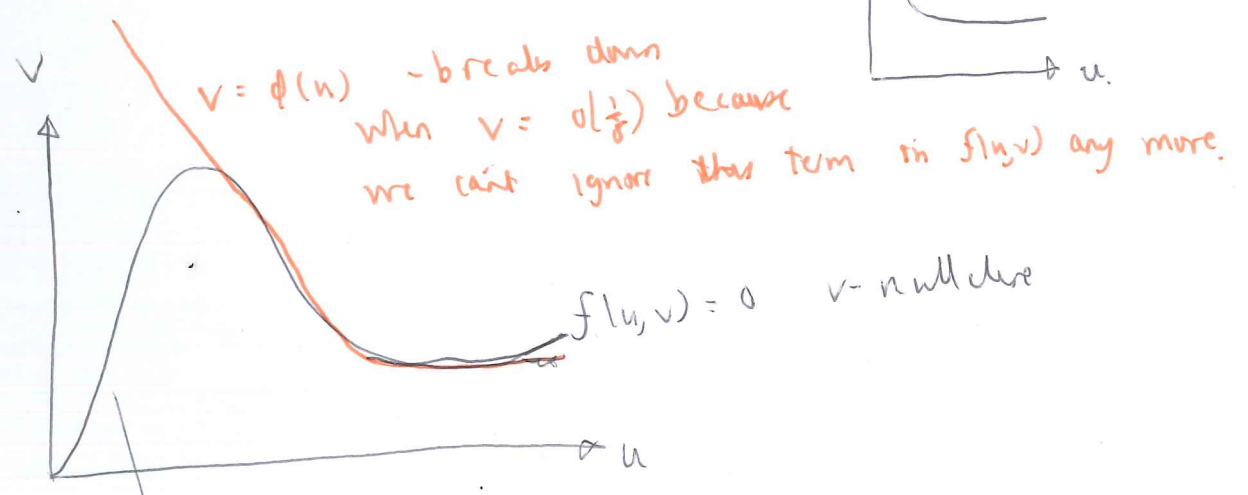
$$\frac{v_m}{1 + v^m} = \frac{\frac{\beta u^n}{1 + u^n}}{\frac{\alpha p + u p}{\alpha p + u p}} \stackrel{\text{def}}{=} \frac{J(u)}{K(u)}$$



$$v = \left[\frac{J(u)}{K(u) - J(u)} \right]^{1/m} = \phi(u)$$



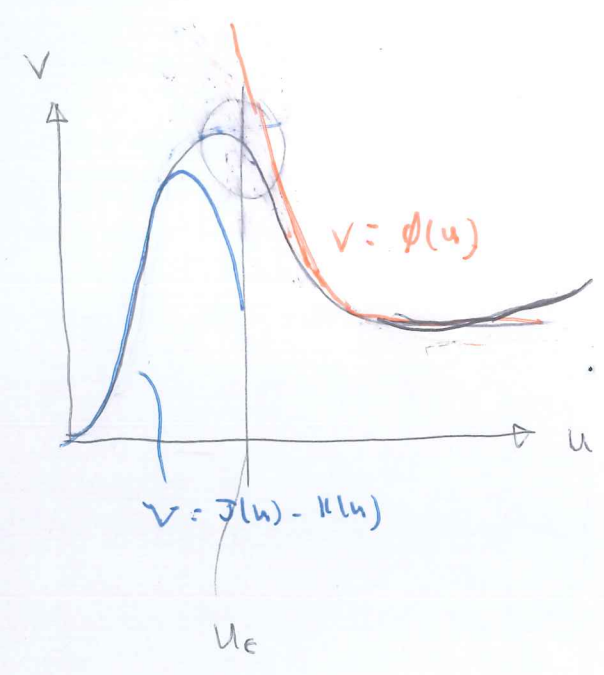
Don't do this in the lecture - this is a problem itself question.



To approximate this part, set $v = \sqrt[m]{\epsilon}$

2) To get the bit when $v = O(\frac{1}{\delta})$, set $v = \frac{V}{\delta}$

Then $f(u, v) = 0 \Rightarrow v = J(u) - K(u)$



Don't do this in the lecture - this is a problem sheet question.

There is a matching region where neither are valid.

Here, we set $u = u_c + \delta^{m/m_1} U$,

$v = \frac{1}{\delta^{1/m_1}} W$

This approximation matches to the inner (blue) and outer (orange) regions. Problem sheet exercise?

Now let's look at the dynamics. The system relaxes
 v rapidly approaches the v -nullcline that we have
 found, ~~but~~ because of the ϵ . ~~But~~ now in the $\frac{dv}{dt}$
 equation. But now if we look at the $\frac{du}{dt}$ equation

we have
$$\frac{du}{dt} = \rho - u - \frac{\gamma}{\epsilon} f(u, v)$$

$$\uparrow$$
 an ϵ here

so we don't just have $u = \text{constant}$ unlike in the
 previous cases. This time we note that

$$\frac{du}{dt} + \gamma \frac{dv}{dt} = \rho - u$$

$$\epsilon \frac{dv}{dt} = f(u, v)$$

in the fast time scale ($t = \epsilon \tau$) we have

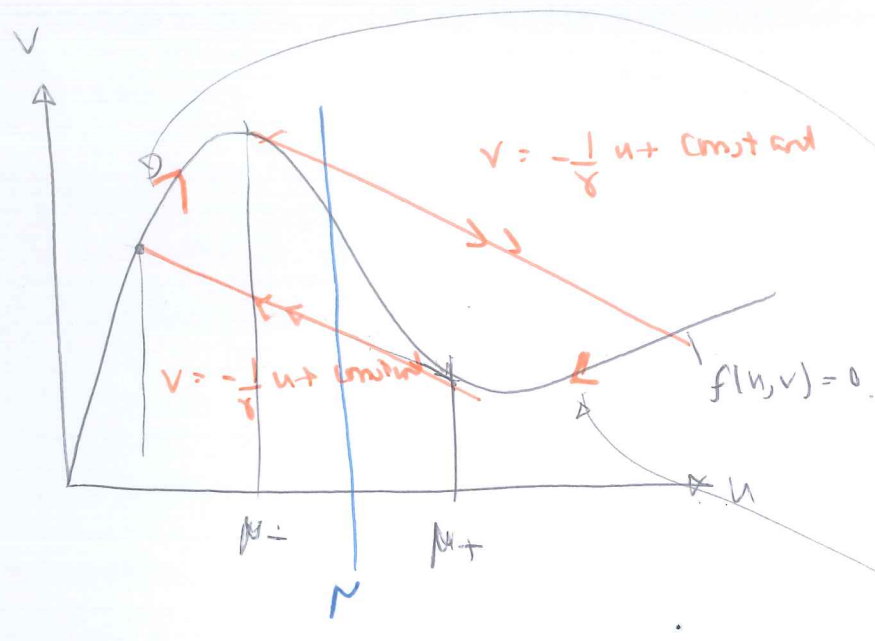
~~$\frac{du}{dt}$~~ $\frac{dv}{d\tau} = f(u, v)$ giving the movement of v to

the v -nullcline and
$$\frac{du}{d\tau} + \gamma \frac{dv}{d\tau} = \epsilon(\rho - u)$$

$$\Rightarrow u + \gamma v = \text{constant}$$

So we move to the v -nullcline along the line

$$v = \frac{\text{constant} - u}{\gamma}$$



μ_+ and μ_- are the places where the curve defined by $f(u, v) = 0$ has gradient $-\frac{1}{\gamma}$

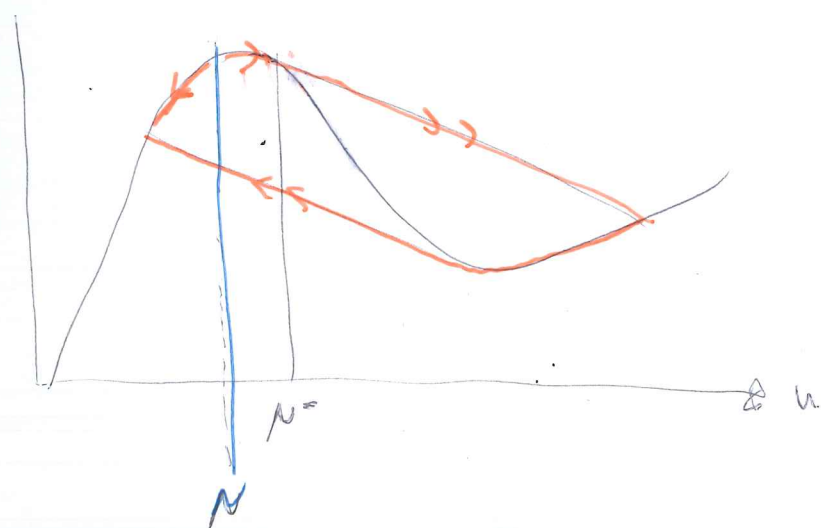
Case (i) $\mu_- < N < \mu_+$

Then $\frac{d}{dt}(u + \gamma v) = \mu_- - u < 0$ when $u > \mu_+$
 > 0 when $u < \mu_-$

This leads to self-sustained or relaxation oscillations

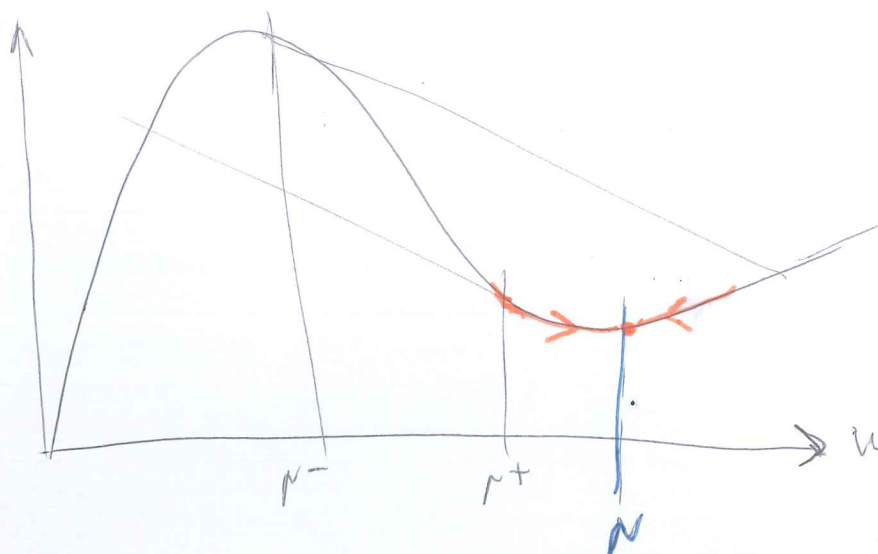
Case (ii) $N < \mu_-$

Then $\frac{d}{dt}(u + \gamma v) = \mu_- - u > 0$ when $u < \mu_-$



Need a bit of excitation to get an excursion - a muscle contraction

Case (iii) $N > N_+$



$$\frac{d}{dt}(u + \gamma v) = N - u$$

Concentration of Ca^{2+} stays high always - Cramp and rigor mortis.

Wave propagation

We have so far seen how an excitable steady state can and spatial variations can lead to travelling solitary waves

(Hodgkin-Huxley)



If you have periodic solutions and spatial variations this can lead to periodic travelling waves. These have been

observed in Xenopus oocytes (sub-sarcomeric frogs) (eggs)

Let's see if we can model this by adding in spatial dependence (diffusion) into our two-pool model:

$$u_t + \gamma v_t = \mu - u + \frac{v u_{xx}}{\epsilon}$$

$$\epsilon v_t = f(u, v)$$

Diffusion of Ca^{2+} in the sarcoplasm

Same model as before plus this extra term

$$v = \frac{D}{l^2 k}$$

Diffusion coefficient of Ca^{2+} in the sarcoplasm

length scale to be determined

leakage rate

Define l is that $v = \epsilon$ to give an interesting (and relevant) balance

This is similar to the Fitzhugh-Nagumo equations with diffusion except this time the diffusion term is in the slow equation rather than the fast one.

Seek travelling wave solutions: $\xi = x + st$ 48

$$u = u(\xi), \quad v = v(\xi), \quad s = \text{wave speed.}$$

$$\Rightarrow s(u' + \gamma v') = \mu - u + \epsilon u'' \quad (1)$$

$$\epsilon s v' = f(u, v) \quad (2)$$

Now we follow the same analysis that we did for Fitzhugh-Nagumo except now we are looking for periodic solutions in ξ rather than a solitary wave.

~~First look at the fast motion by re~~
As before, (2) tells us that we quickly move onto the v nullcline $f(u, v) = 0$.
Rescale $\xi = \epsilon X$ to see how we get there. This gives

$$\begin{aligned} s(u' + \gamma v') &= u'' \quad (3) & \left(' = \frac{d}{dX} \right) \\ s v' &= f(u, v) \quad (4) & \text{(to leading order in } \epsilon) \end{aligned}$$

Integrate (3) $\Rightarrow u' = s [u - u_0 + \gamma(v - v_0)]$

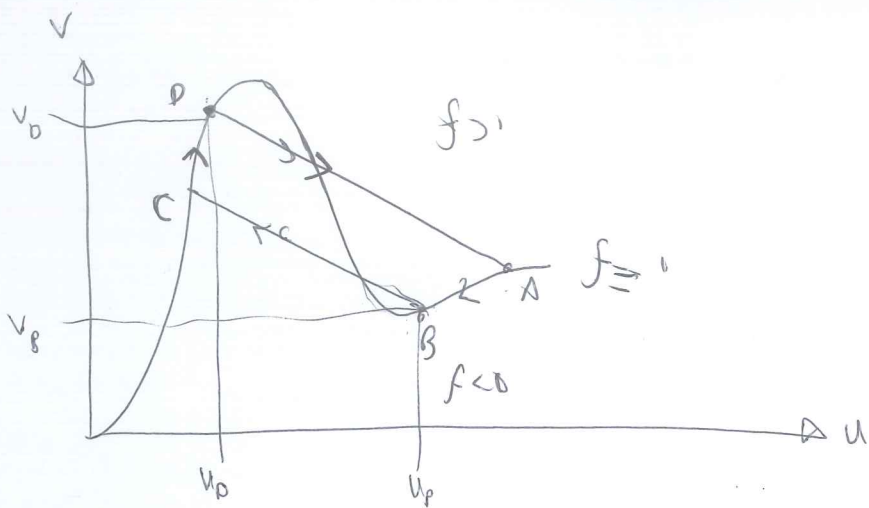
Where $u = u_0, v = v_0$
at the place where $u' = 0$
call this point D .

Now let's shift the origin to (u_0, v_0) :

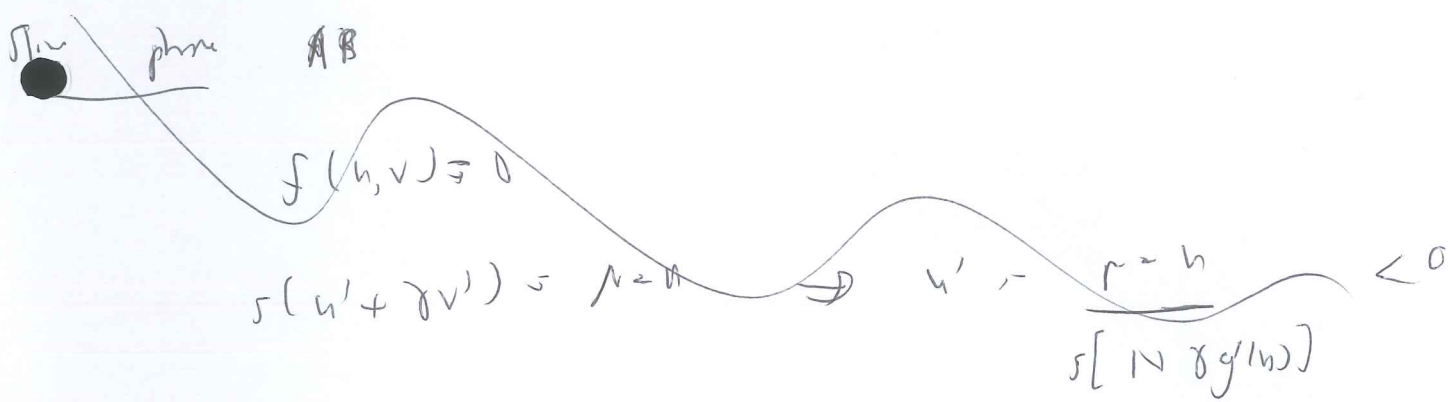
$$u = u_0 + \tilde{u}, \quad v = v_0 + \tilde{v}$$

Then we have

$$\begin{aligned} \tilde{u}' &= s [\tilde{u} + \gamma \tilde{v}] \\ s \tilde{v}' &= f(\tilde{u}, \tilde{v}) \end{aligned}$$



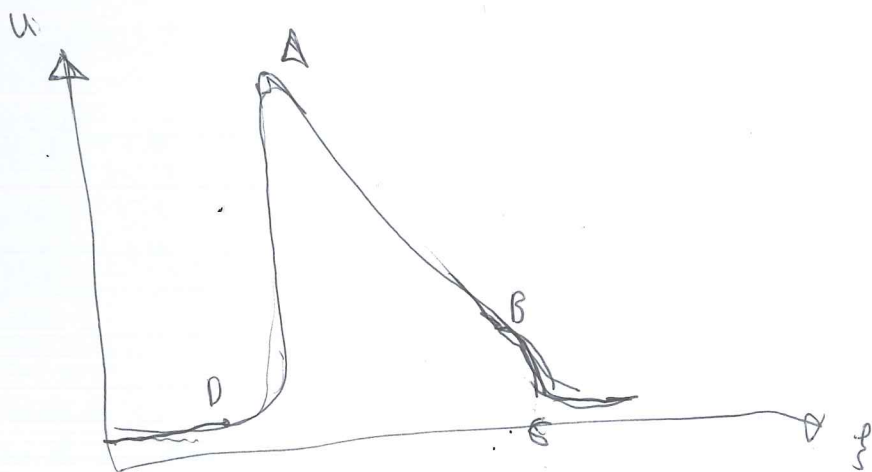
Hypothesized
 phase plane
 We will prove
 why it looks
 like this but
 good to know
 what we are
 looking for.



the jumps at points

We don't know where (u_0, v_0) and (u_p, v_p) are and
 need to find them as part of the solution.

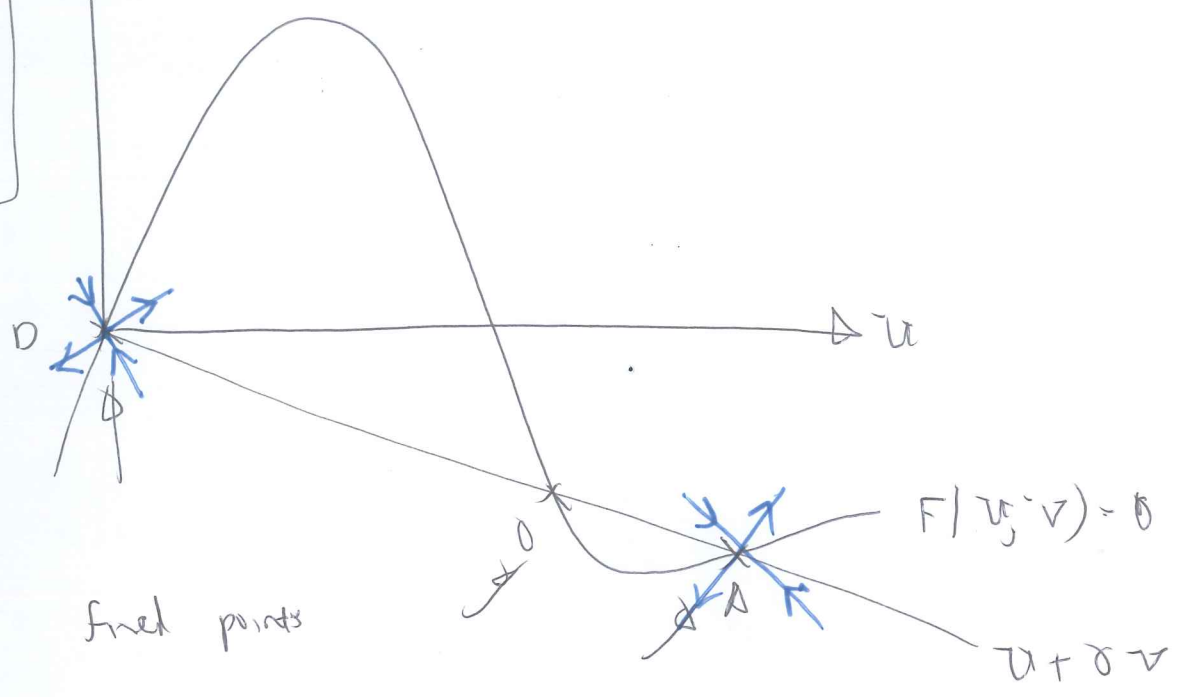
Hypothesized oscillated wave (which repeats periodically)



D and A are fixed points of this u, v system

$$u' = s[-u + \delta v]$$

$$s-v = f(u, v)$$



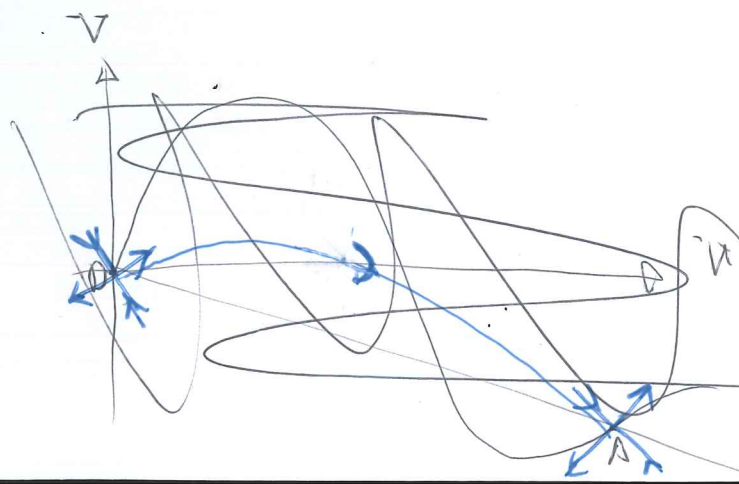
fixed points

Fixed points: $u + \delta v = 0$
 $f(u, v) = 0$

A and D are saddles
 O is an unstable node or focus

Can be shown using standard linear stability analysis

Next step is to look up D and A:

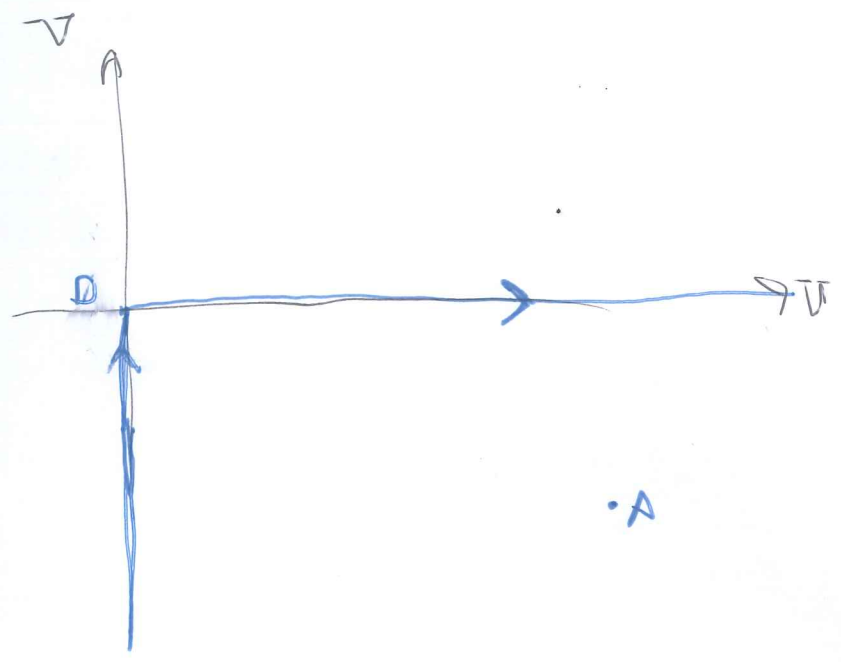


There is a unique value of s that links up D + A (see next page for proof)

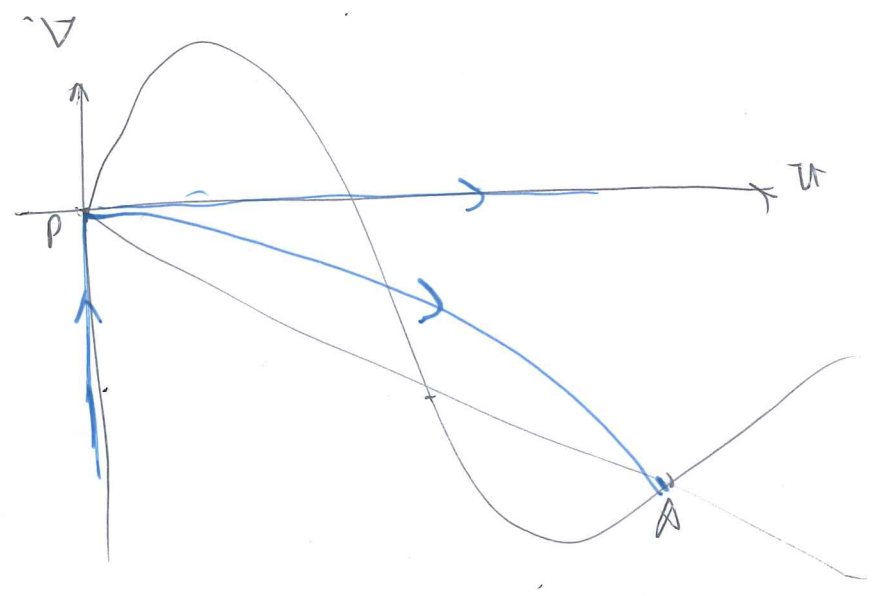
There is a unique value of s that joins D to A 51

You can see this because if ~~$s \rightarrow 1$~~ $s \rightarrow 1$ and $V < 0$
then $U = \text{constant}$ and V goes up to D

And if $s \rightarrow \infty$ then $U \rightarrow \infty$



\therefore there must be something that joins D to A for an intermediate value of s



Now we go from A to B along ~~the~~ $f(u,v) = 0$ on the v -nullcline. This is the slow part:

$$s(u' + \delta v') = \mu - u + \epsilon u'' \implies s(u' + \delta v') = \mu - u$$

$$\epsilon \delta v' = f(u,v)$$

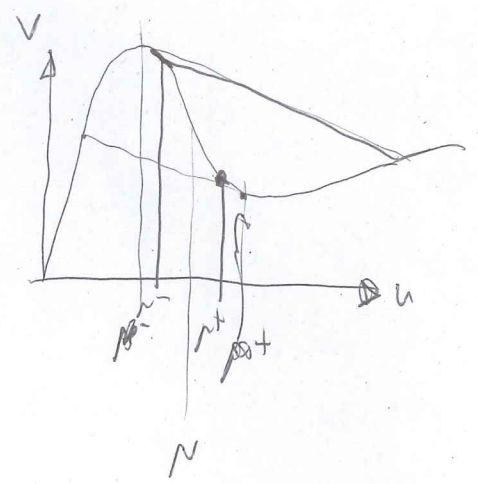
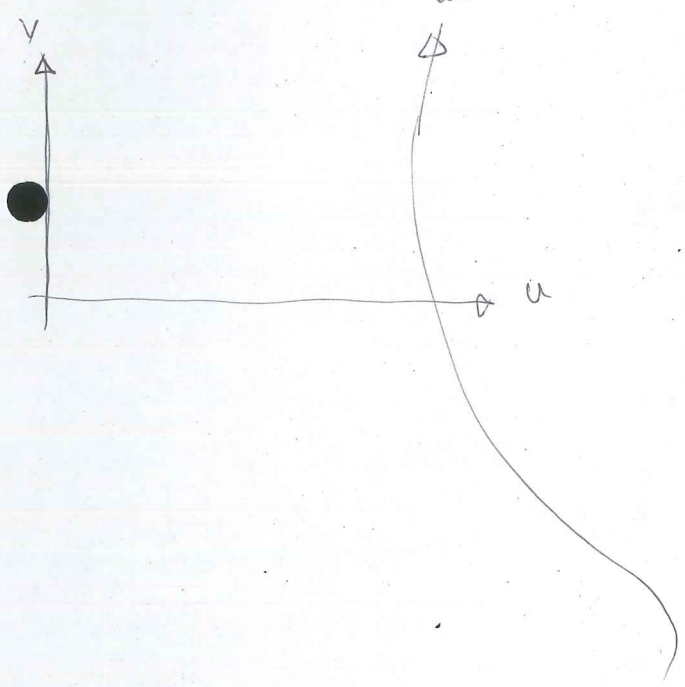
$f(u,v) = 0$
to leading order in ϵ

We know that B is to the left of A because we can show that $u' < 0$ on this slow phase.

Proof

$$u' = \frac{\mu - u}{s(1 + \delta \frac{v'}{u'})}$$

$$= \frac{dv}{du}$$



will
We still assume $\mu_- < N < \mu_+$
so $\mu - u < 0$

Recall μ^-, μ^+ are the places where the gradient of the curve $f(u,v) = 0$ equals $-\frac{1}{\delta}$

so we know this is $> -\frac{1}{\delta}$

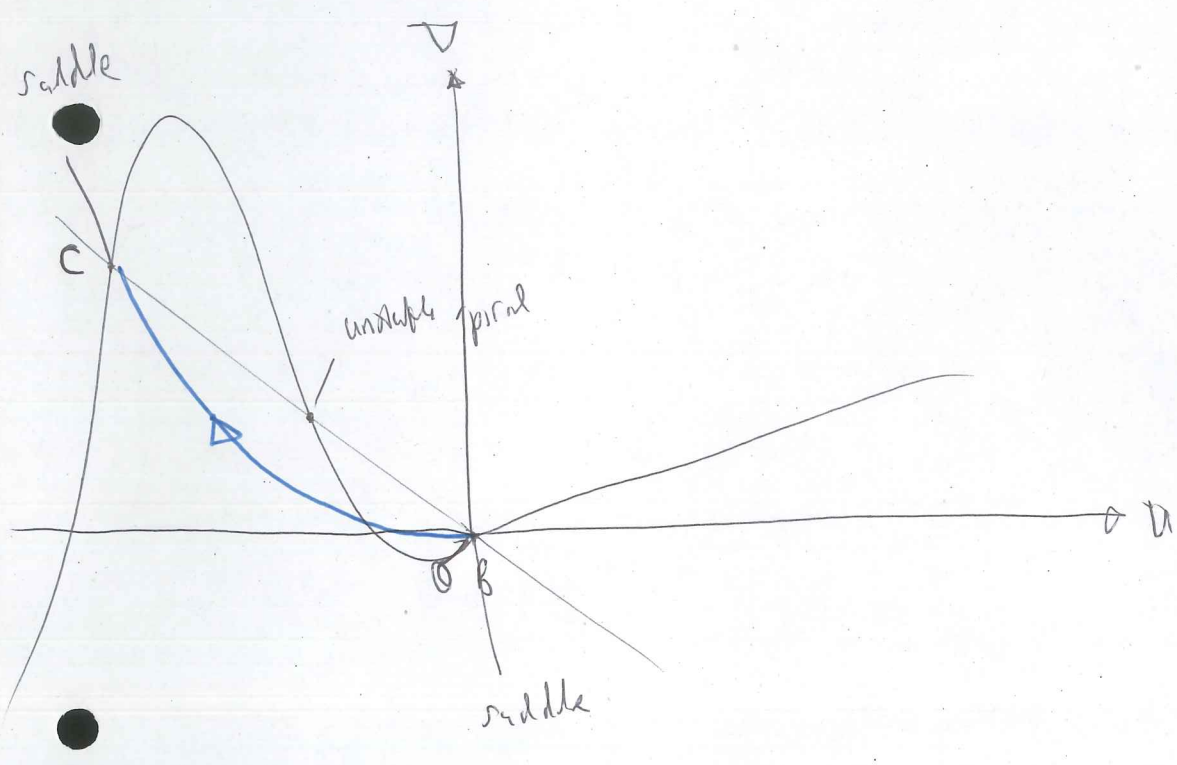
so denominator $> 0 \implies u' < 0 \implies$ trajectory moves down $f(u,v) = 0$ to the left.

Now do we know where B is? The $F(u, v) = 0$ curve (unique) -
 place where we link back on a fast trajectory

To analyze this, we set $u = u_B + \tilde{u}$,
 $v = v_B + \tilde{v}$

to get (like before) $\tilde{u}' = s(\tilde{u} + \gamma \tilde{v})$
 $s \tilde{v}' = F(\tilde{u}, \tilde{v})$

Fixed points:
 $\tilde{u} + \gamma \tilde{v} = 0$
 $F(\tilde{u}, \tilde{v}) = 0$



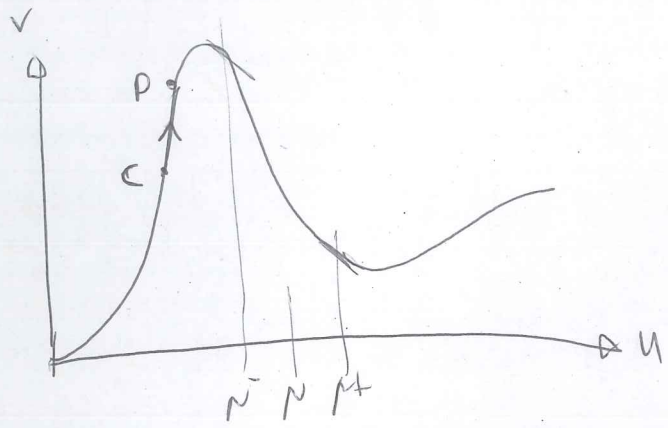
B and C are fixed points of this (u, v) system

The values of u_B, v_B are set so that we link back up to the other fixed point C, in this new (u, v) phase plane (note this one is different to the previous (u, v) phase plane because now we are centered around (u_B, v_B) rather than (u_0, v_0)).

Thus then leaves the final slow phase C + D.

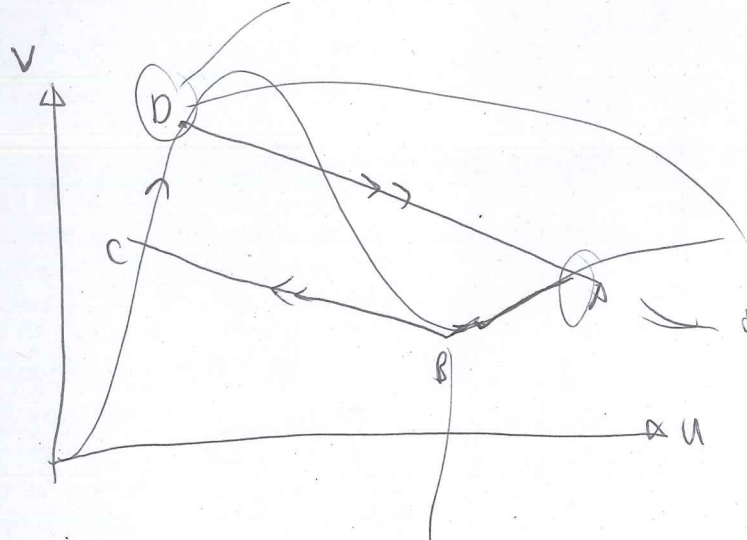
As on pt. C we have $u' = \frac{r-u}{s \left[1 + \gamma \frac{dv}{du} \right]} > 0$ as $u < r$

so this takes u to the right on the curve $f(u, v) = 1$



~~NA that although~~
 so what we have:

~~set by fixed point (u, v) system~~



fixed
 set by fixed points (u, v)
 system; s chosen
 to link the two
 trajectories

set by fixed points (u, v)
~~fixed~~ second (u, v) system;
 (u_B, v_B) chosen to link the two
 trajectories

Note that although u_B (and thus v_B) were fixed,
 u_D (and thus v_D) were not. So there is a one-parameter
 family of periodic travelling waves, with corresponding
 wave speeds.

The heart

58

There are two parts to the heart function:

- 1) Electrochemical action - causes muscle contraction to pump blood around the body.
- 2) Mechanical action - enables unidirectional circulation via a system of valves.

We must begin by studying 1 and then move onto 2

The electrochemical action of the heart

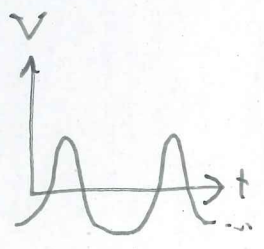
The heart has four chambers



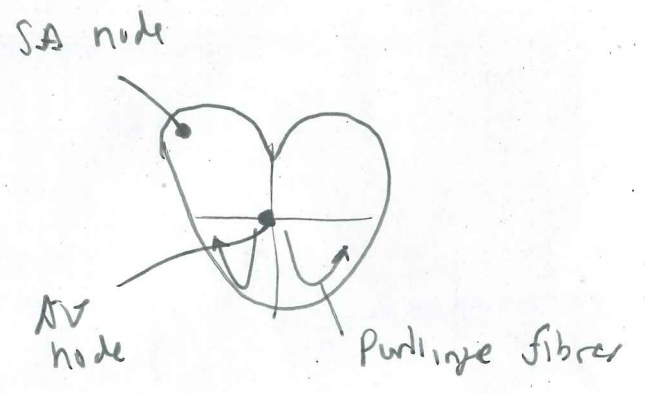
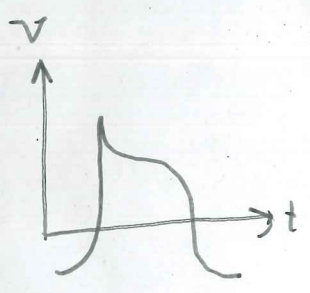
Blood flows into the RA from the venous system, to the RV, perfuses through the lungs (gains oxygen), moves to the LA then the LV then to the arteries

- RA = right atrium
- LA = left atrium
- RV = right ventricle
- LV = left ventricle

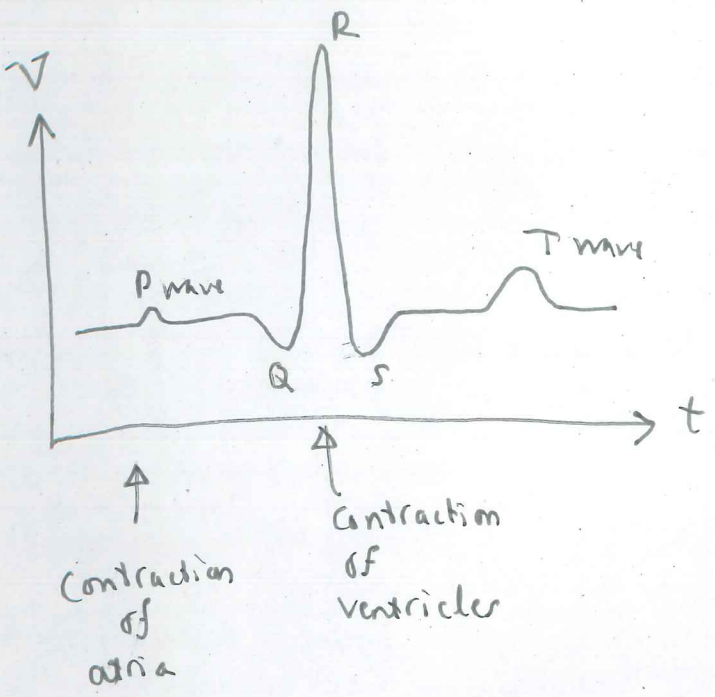
In the RA is the sino-atrial (SA) node, whose cells act as pacemakers with a periodic action potential



Other cells (atrial/ventricular ^(AV) myocytes, AV node, Purkinje fibres) are excitable with a distinct action potential.



The electrocardiogram (ECG)



P = depolarisation of the atria
 Q, R, S = depolarisation of the ventricles
 T = repolarisation of the ventricles.

Approximately 20 waves propagate through the heart from SA. We will learn about these

misregulation of conduction paths can lead to 're-entrant' spiral waves, which cycle round the diseased tissue. This causes ventricular tachycardia.

the diseased heart, spiral waves can become chaotic. This causes ventricular fibrillation.

Noble model (1982)

was an early model for the action potential of ventricular myocytes similar to Hodgkin-Nuxley but more variables due to greater number of currents involved.

Represents the ionic current as the sum of sodium, potassium and leakage currents:

$$C_m \frac{dV}{dt} = -I_i \quad I_i = I_{Na} + I_K + I_L$$

residual Hodgkin-Huxley

$$I_{Na} = [g_0 + g_{Na} m^3 h] (V - V_{Na})$$

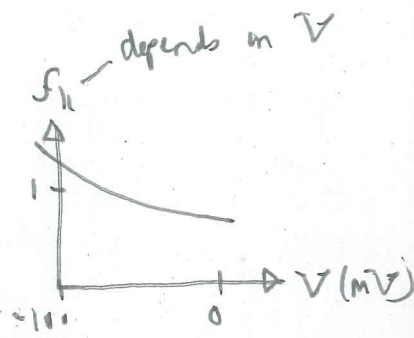
switching on gate switching off gate

conductivity of Na (m and h gates)

$$I_K = [f_{K1} + g_{K2} n^4] (V - V_K)$$

instant long lasting

conductivity of K (n gate only)



$$I_L = g_L (V - V_L)$$

constant leakage conductivity.

ex. this

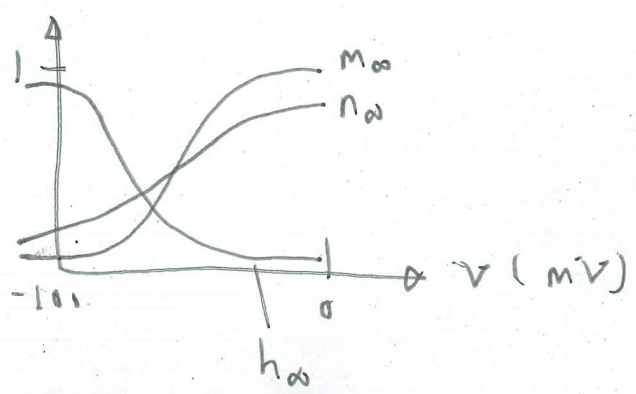
All of this comes largely from experimental fitting but the key idea is the gates

and the gate variables satisfy:

$$\tau_m \dot{m} = m_\infty - m \quad \tau_m \sim 0.25 \text{ ms}$$

$$\tau_h \dot{h} = h_\infty - h \quad \tau_h \sim 8 \text{ ms}$$

$$\tau_n \dot{n} = n_\infty - n \quad \tau_n \sim 50 \text{ ms} - \text{much longer.}$$



We non-dimensionalize via

$$V = \underbrace{(V_k)}_{(V_k < 0)} \hat{V}, \quad t = T_h \hat{t}$$

to get

$$\left(\begin{array}{c} T_m \\ T_n \end{array} \right) \dot{m} = m_a - m \quad \Rightarrow \quad m \approx m_a \quad (\text{rapidly reached})$$

$$h = h_a - h$$

$$\left(\begin{array}{c} T_n \\ T_m \end{array} \right) \dot{n} = n_a - n \quad \dot{n} = \varepsilon (n_a - n)$$

$\gg 1 = \frac{1}{\varepsilon}$, say
(= 0.016)

$$\dot{V} = -G(V, h, n)$$

This is the dimensionless form
of $C_m \dot{V} = -J_i$ from previous page

(dropping hats on dimensionless V)

with $G = - \left[\gamma_0 + \gamma_{Na} M_a^3 (V) h \right] (V_{Na} - V)$

$$\left(\begin{array}{c} \ll 1 \\ \gg 1 \end{array} \right) + \phi(V+1) + \gamma_L (V + V_L)$$

we will use this later.

$$= \phi_{||}(V) + \gamma_{||} n^4$$

we will use this later

Note took $\gamma_L = 0$ to begin with (this is the leakage part)

First consider the fast phase. (ie no time rescalings)

This gives $n = \text{constant}$ to leading order in ϵ .

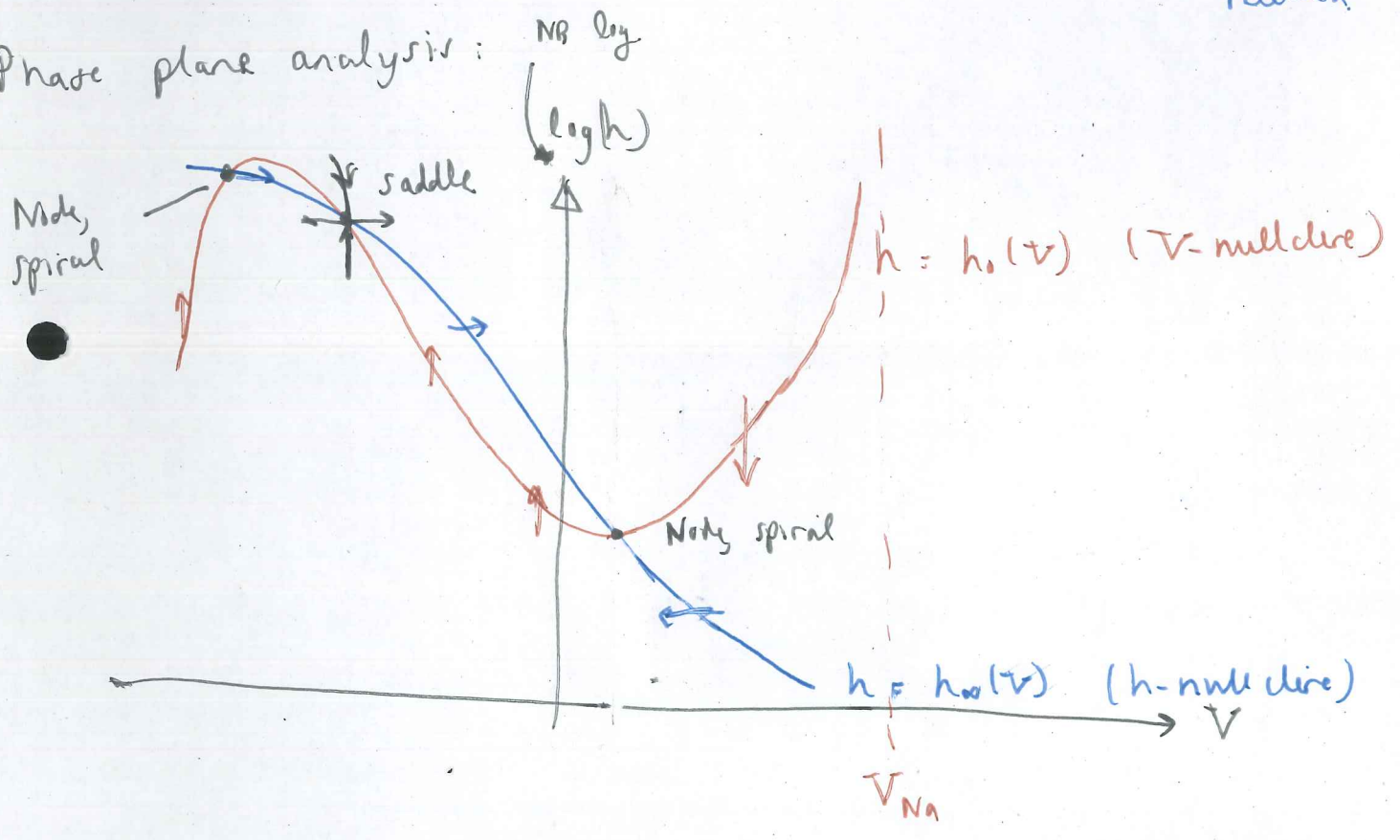
$$\dot{h} = h_{na} - h$$

$$\dot{V} = -\gamma_{Na} m_0(V)^2 [h_0(V) - h] (V_{Na} - V)$$

where $h_0(V) \stackrel{\text{def}}{=} \frac{1}{\gamma_{Na} m_0(V)^2} \left(\frac{\rho(V+1) + \gamma_L(V+V_L) - \gamma}{V_{Na} - V} \right)$

$\stackrel{\text{def}}{=} s$ NA sure if this is needed

Phase plane analysis:



We need to find the stability of the two nodes

Linearising about these fixed points (h^*, v^*) , via

$h = h^* + N, \quad v = v^* + w$ gives

$$\frac{d}{dt} \begin{pmatrix} H \\ W \end{pmatrix} = \underbrace{\begin{pmatrix} -1 & h_0' \\ A & -\Delta h_0' \end{pmatrix}}_{\underline{M}} \begin{pmatrix} H \\ W \end{pmatrix}$$

Maybe we don't need to do this in lectures and we can just say they are stable

where $A = \delta_{Na} m_a^3(v)(v_{Na} - v)$

$\text{tr}(\underline{M}) = -1 - \Delta h_0'$

Left fixed point

Here, $h_0' > 0$ and m_a is quite small for low v values (see colour graph) so $m_a \ll 1$ so $A \ll 1$.

So $\text{tr}(\underline{M}) < 0 \Rightarrow$ stable

Right fixed point

$h_0' < 0$ so $\text{tr}(\underline{M}) < 0$ so stable too.

Now we need to consider the slow variations in n .

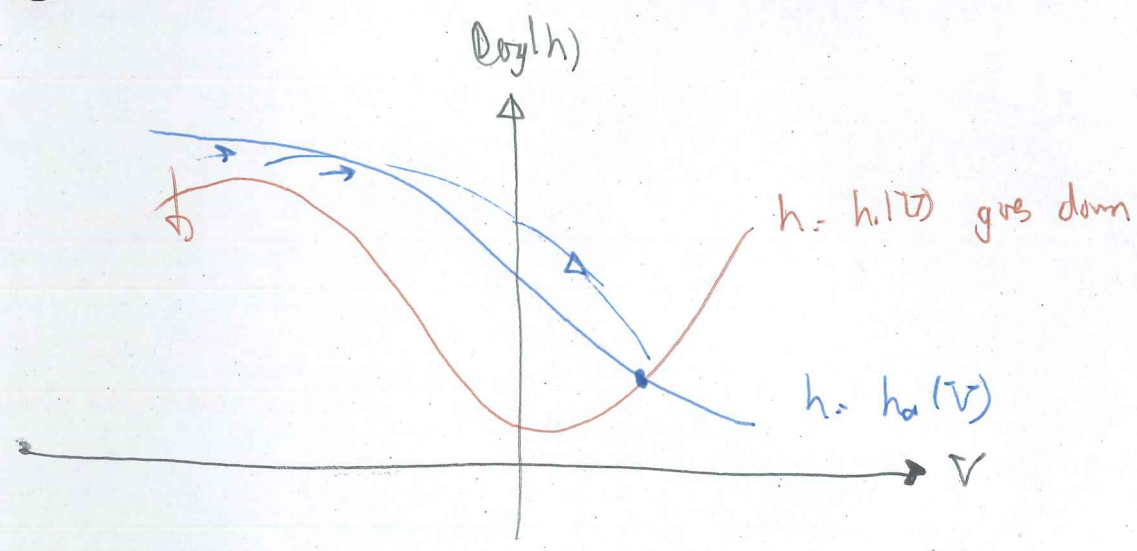
$$\dot{n} = \epsilon (n_{\infty} - n)$$

When solution is at left fixed point, V is low so n_{∞} is low (from previous graph) so $n_{\infty} - n < 0$ so n goes down

so ϕ goes down (recall $\phi = \bar{\phi}_u(V) + \delta_u \underline{n}$) so

$h_1(V)$ goes down (see definition of $h_1(V)$ so phase (and actually, changes shape)

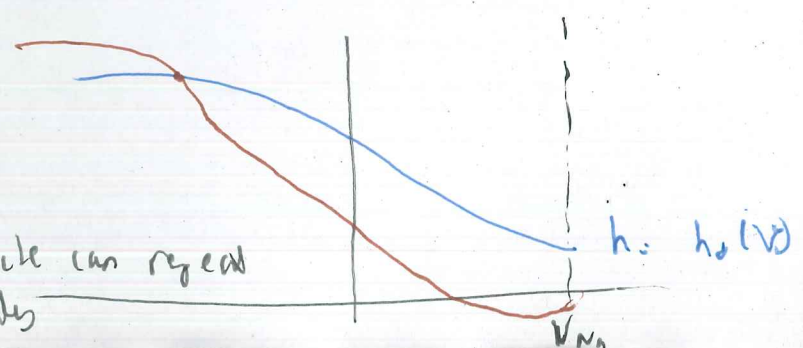
phase evolves to:



So this is something different because in this phase plane the nullclines are moving

In doing so the left fixed point disappears and so the system jumps to the right fixed point

From here, n_{∞} is now high so $n_{\infty} - n > 0$ so n goes up so the $h = h_0(V)$ curve moves up until we get a loss of the right fixed point:



The left fixed point reappears as $h = h_0(V)$ graph also changes shape.

This cycle can repeat periodically

Waves in two or three dimensions

Waves propagate through the heart in approximately two dimensions

Periodic wave propagation

Let \underline{w} be a vector of 'reactants' (eg v, m, n, h) with

kinetics $\frac{\partial \underline{w}}{\partial t} = f(\underline{w})$

Suppose \underline{w} has a stable limit cycle behaviour $\underline{w} = \underline{w}_0(t)$ with period T .

Now consider weak diffusion to add spatial dependence:

$$\frac{\partial \underline{w}}{\partial t} = f(\underline{w}) + \varepsilon \nabla^2 \underline{w} \quad (1) \quad \varepsilon \ll 1.$$

(This assumes all reactants have the same diffusion coefficient.)

This weak coupling causes \underline{w} to evolve on a slow time scale, $\tau = \varepsilon t$.

We seek a multiple scales approximation: $\underline{w} = \underline{w}(\underline{x}, t, \tau)$.

Then (1) becomes $\frac{\partial \underline{w}}{\partial t} + \varepsilon \frac{\partial \underline{w}}{\partial \tau} = f(\underline{w}) + \varepsilon \nabla^2 \underline{w}$ (2)

We seek a series approximation $\underline{w} = \underline{w}_0 + \varepsilon \underline{w}_1 + \dots$

At $O(1)$ in ② this gives

$$\frac{\partial \underline{w}_0}{\partial \tau} = f(\underline{w}_0)$$

$$\Rightarrow \underline{w}_0 = \underline{w}_0(t + \psi(x, \tau))$$

↑

This tells us how the phase can shift in space and on the long time τ .

(Recall \underline{w}_0 is the solution to $\frac{\partial \underline{w}}{\partial \tau} = f(\underline{w})$ so the phase shift is like the constant of integration and the only way to get the x and τ dependence in.)

At $O(\varepsilon)$ we get

$$\frac{\partial \underline{w}_1}{\partial \tau} - J \underline{w}_1 = - \left[\frac{\partial \psi}{\partial \tau} - \nabla^2 \psi \right] \underline{w}_0' + |\nabla \psi|^2 \underline{w}_0''$$

$$\text{where } J = Df(\underline{w}_0(t))$$

↑
Derivative with respect to w_0 .

This is a linear inhomogeneous equation.

We also note that $\underline{s} \stackrel{\text{def}}{=} \underline{w}_0'$ satisfies

$$\frac{\partial \underline{s}}{\partial \tau} - J \underline{s} = 0$$

$$\left(\begin{array}{l} \text{since } \frac{\partial \underline{w}_0}{\partial \tau} = f(\underline{w}_0) \\ \text{" } \frac{\partial}{\partial \tau} (\underline{w}_0') = \frac{df}{dw} \underline{w}_0' \\ \quad \quad \quad \downarrow \\ \quad \quad \quad = J \underline{s} \end{array} \right)$$

Thus we may use the Fredholm alternative to determine the solution for \underline{w}_1 , which gives

$$\underline{w}_1 = -t \left[\frac{\partial \psi}{\partial t} - \nabla^2 \psi \right] \underline{\xi} + |\nabla \psi|^2 \underline{u} \quad \text{as } t \rightarrow \infty$$

where $\underline{u} = \underline{\xi} \left[\bar{\alpha} t + p(t) \right]$
 $\bar{\alpha}$ a parameter that depends on $\underline{\xi}$
 $p(t)$ a periodic function

(No need to know the details of this.)

Now we choose ψ to remove the secular terms that grow unboundedly with time (underlined in orange)

Thus we choose

$$\frac{\partial \psi}{\partial t} = \nabla^2 \psi + \bar{\alpha} |\nabla \psi|^2$$

This is an integrated version of Burgers' equation - in one dimension, $u = -\frac{\partial \psi}{\partial x}$ satisfies $\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} = \frac{\partial^2 u}{\partial x^2}$

(Burgers' equation).

Solutions of Burgers' equation include shocks.

Target patterns

Consider the equation $\frac{\partial \psi}{\partial \tau} = \nabla^2 \psi + \bar{\alpha} |\nabla \psi|^2$

with boundary conditions $\psi = \Omega \tau$ at $r = b$

(this could correspond to an impurity)

↑
radial coordinate

This corresponds to patterns that are formed by a pacemaker in the heart in the SA node and a radiation condition $\frac{\partial \psi}{\partial r} < 0$ as $r \rightarrow \infty$ - this generates circular patterns that originate from a point. (This ensures that the waves move outwards.)

We make the Ansatz $\psi = \Omega \tau - f(r)$

This then gives

$$f'' + \frac{1}{r} f' - \bar{\alpha} (f')^2 + \Omega = 0$$

with $f(b) = 0$

and $f' \rightarrow 0$ as $r \rightarrow \infty$

The solutions to this are

$$f(r) = -\frac{1}{\bar{\alpha}} \log \left[\frac{K_0(\sqrt{\bar{\alpha} \Omega} r)}{K_0(\sqrt{\bar{\alpha} \Omega} b)} \right]$$

K_0 = modified Bessel function of the second kind of order zero

K_0 is the solution to

$$g'' + \frac{1}{r} g' - g = 0$$

that decays as $r \rightarrow \infty$ (The other solution which $\rightarrow \infty$ as $r \rightarrow \infty$ is J_0)

Spiral waves

Now consider the ansatz $\psi = \Omega T + m\theta - g(r)$ $m \in \mathbb{N}$

with boundary condition $\psi = \Omega T + m\theta$ at $r = b$

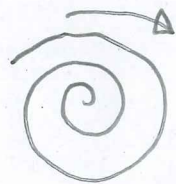
and the radiation condition again, $\frac{\partial \psi}{\partial r} < 0$ as $r \rightarrow \infty$.

This time, the solution is

$$g = -\frac{1}{\alpha} \log \left[\frac{K_0(\sqrt{\alpha \Omega} r)}{K_0(\sqrt{\alpha \Omega} b)} \right] \quad v = i\bar{\omega} m$$

and $\psi \sim \Omega T + m\theta - \sqrt{\frac{\Omega}{\alpha}} r$ as $r \rightarrow \infty$

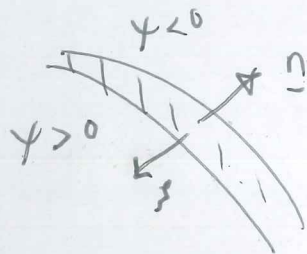
This is an Archimedean spiral



This spiral rotates and continues to excite the tissue.

Curved front propagation

Consider a wave in which the phase ψ of the potential vane rapidly within a thin region (the wave front) which curve more slowly in other directions



● Suppose that the equation $v_t = f(v) + \nabla^2 v$ (*) has a one-dimensional travelling wave solution $v = v(\xi)$, $\xi = ct - x$, $c > 0$, $v(\infty) = 0$, $v(-\infty) = 0$

$$\Rightarrow cv' = f(v) + v'' \quad (†)$$

● ψ denote the phase of the wave. We define the location of the wavefront as $\psi(x, t) = 0$ (eg $\psi = ct - x$ is a target pattern)

Let s be the distance along the normal \underline{n} in the inward direction (see above sketch) $\underline{n} = -\frac{\nabla\psi}{|\nabla\psi|}$ by

definition of the normal to a surface ψ .

$$\begin{aligned} \text{So } \delta\psi &= \nabla\psi \cdot \delta\mathbf{x} \\ &= -|\nabla\psi| \underline{n} \cdot \delta\mathbf{x} \\ &= -|\nabla\psi| \delta s \end{aligned} \Rightarrow \psi_s = |\nabla\psi|$$

We look for a solution of the form $v(\psi)$ 70

In (*) this gives

$$v'(\psi) (\psi_t - \nabla^2 \psi) = f(v) + v''(\psi) |\nabla \psi|^2$$

Since the front is thin this allows us to make a quasi one dimensional approximation: $v_t \approx |\nabla \psi| v'(\psi)$

$$\Rightarrow v_{\psi\psi} + \frac{v_{\psi}}{|\nabla \psi|} \left[\nabla^2 \psi - \frac{\partial}{\partial t} |\nabla \psi| - \psi_t \right] + f(v) = 0$$

Comparing this with (*) which holds if we have a travelling wave solution, we see that we need

$$\frac{1}{|\nabla \psi|} \left[\nabla^2 \psi - \frac{\partial}{\partial t} |\nabla \psi| - \psi_t \right] = -c$$

$$\Rightarrow \psi_t = \nabla^2 \psi - \frac{\partial}{\partial t} |\nabla \psi| + c |\nabla \psi|$$

$$\Rightarrow \frac{\psi_t}{|\nabla \psi|} = c - \underbrace{\nabla \cdot \left(\frac{-\nabla \psi}{|\nabla \psi|} \right)}_{\underline{n}} \quad (*)$$

But now $\psi = 0$ denotes the front so $\frac{D\psi}{Dt} = 0$

$$\Rightarrow \psi_t + \underline{v} \cdot \nabla \psi = 0$$

$$\Rightarrow \psi_t + v_n |\nabla \psi| = 0$$

$$\Rightarrow v_n = - \frac{\psi_t}{|\nabla \psi|}$$

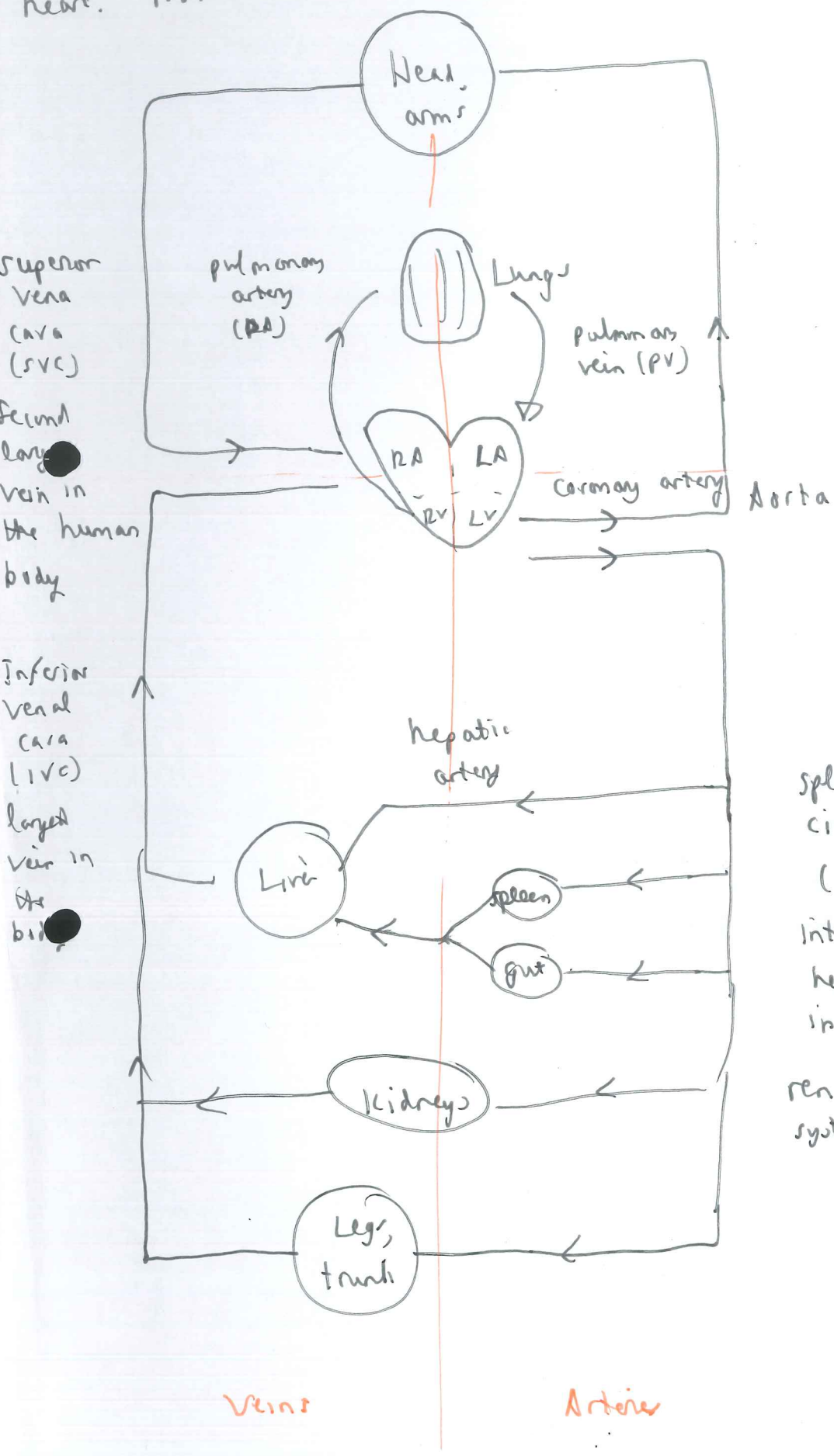
So in (*) \Rightarrow

$$v_n = c - \nabla \cdot \underline{n}$$

This is the Eikonal equation

The heart as a pump

We have so far looked at the electrochemical action of the heart. Now we will move on to look at the mechanical action.

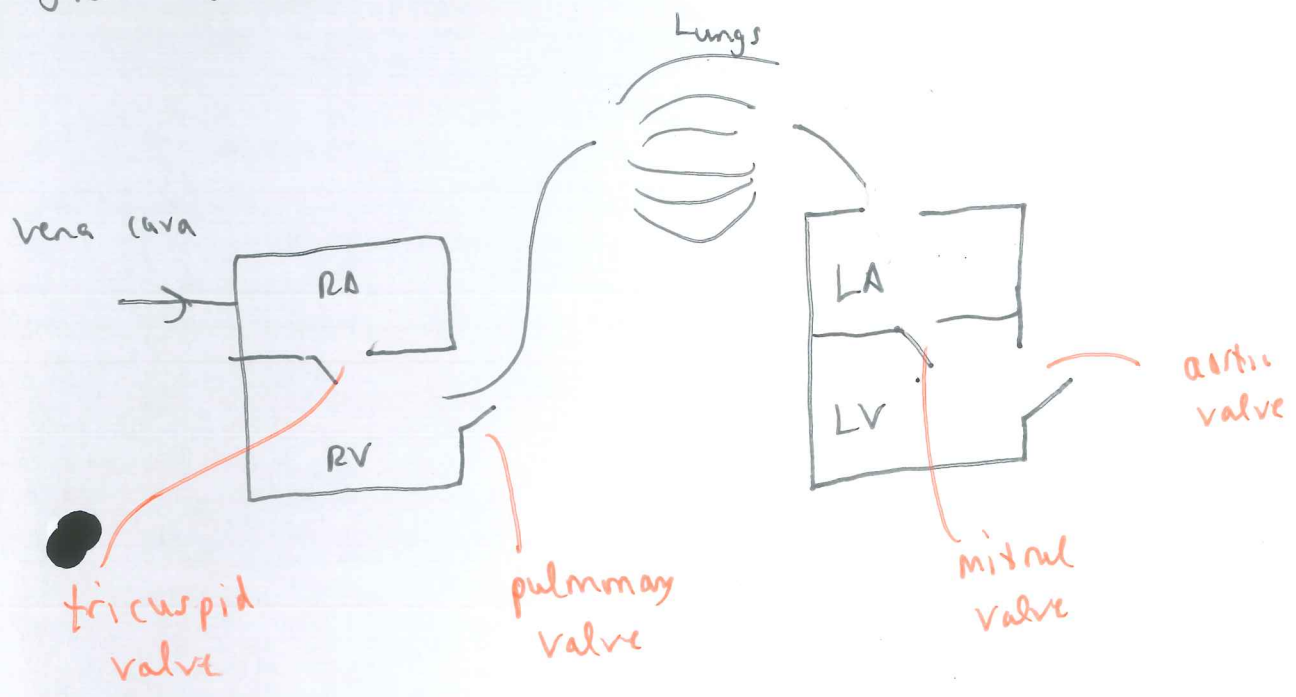


RA = right atrium
 RV = right ventricle
 LA = left atrium
 LV = left ventricle

splanchnic circulation
 (gastro, small
 intestinal, colon, pancreas,
 hepatic and spleen circulation)
 in parallel with one another.

renal system

Blood collects oxygen (O_2) from the lungs and delivers this to the tissues and collects CO_2 from tissues and dumps this to the lungs (to exhale)



We have looked at

~~There are two parts to the heart function:~~

- 1) electrochemical action - causes muscle contraction to pump blood around the body.

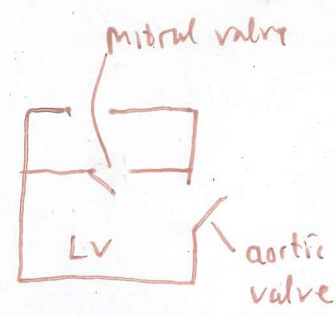
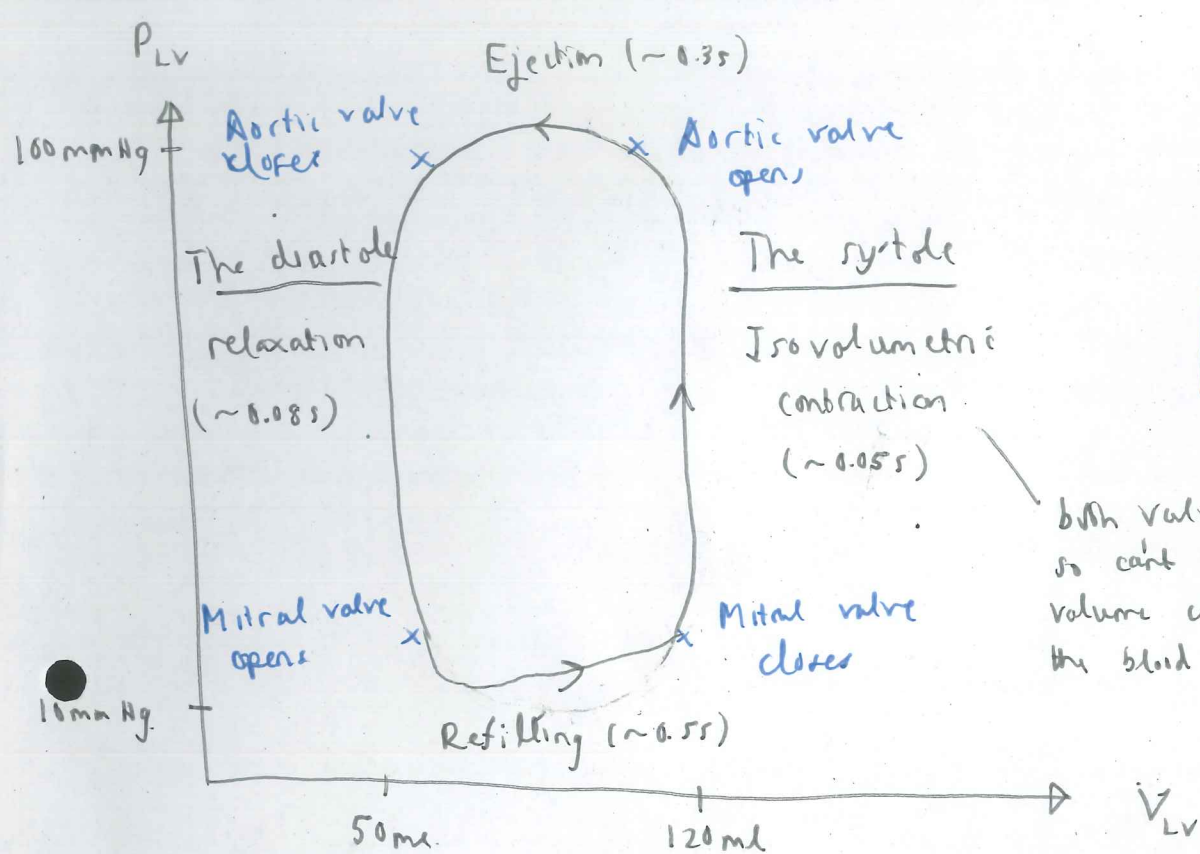
Now we will look at

- 2) mechanical action - enables unidirectional circulation via a system of valves.

~~We will begin by studying 1 and then move onto 2.~~

Pressure-volume cycle of the left ventricle

(The right atr, almost synchronously)



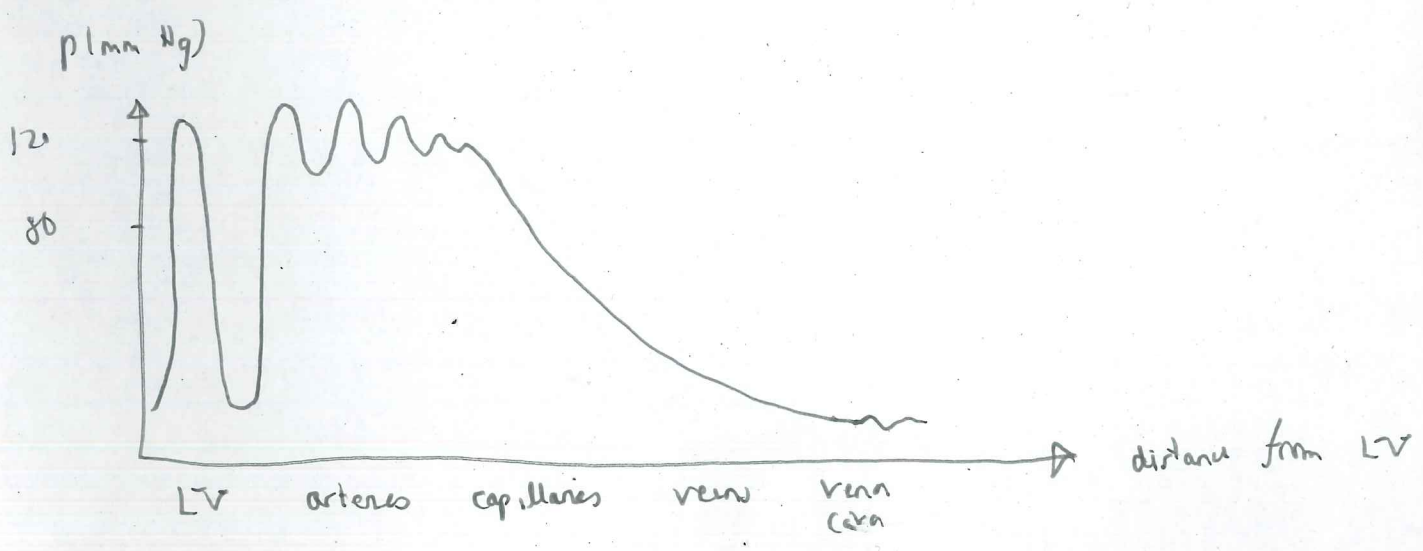
both valves closed so can't have any volume change because the blood is incompressible

stroke volume = change of left ventricular volume on contraction = 70 ml

Systole = contraction phase - ventricular pressure rises
 Diastole = relaxation phase - ventricular pressure falls

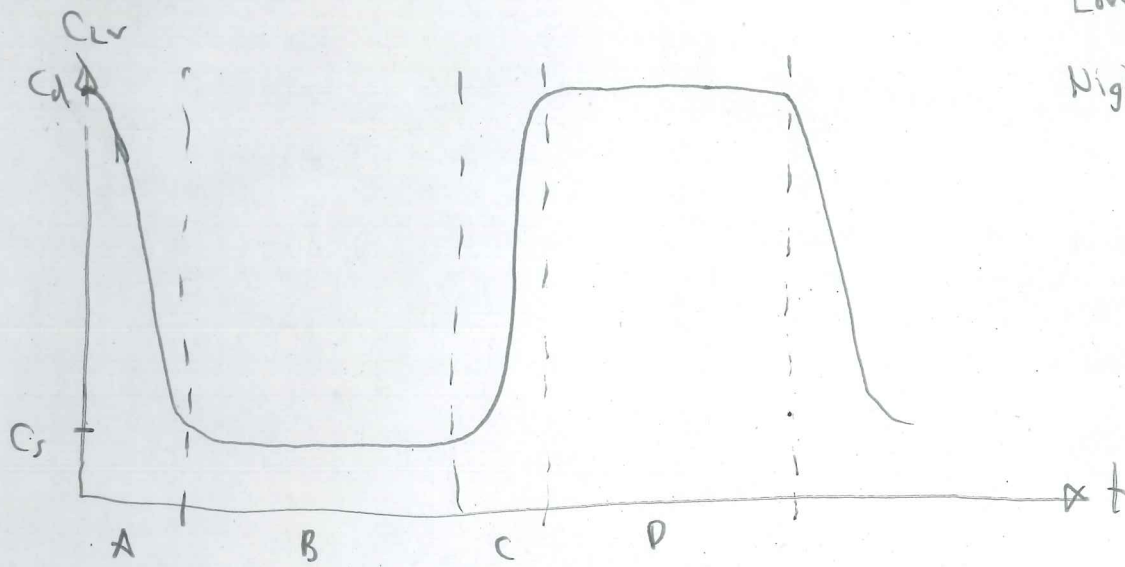
Heart rate = $\frac{1}{\text{period of the sino atrial node cells}}$

[Do next page before the picture]



The compliance can be thought of as a prescribed periodic function: $C_{LV} = \frac{1}{\text{elastance}}$ of the heart

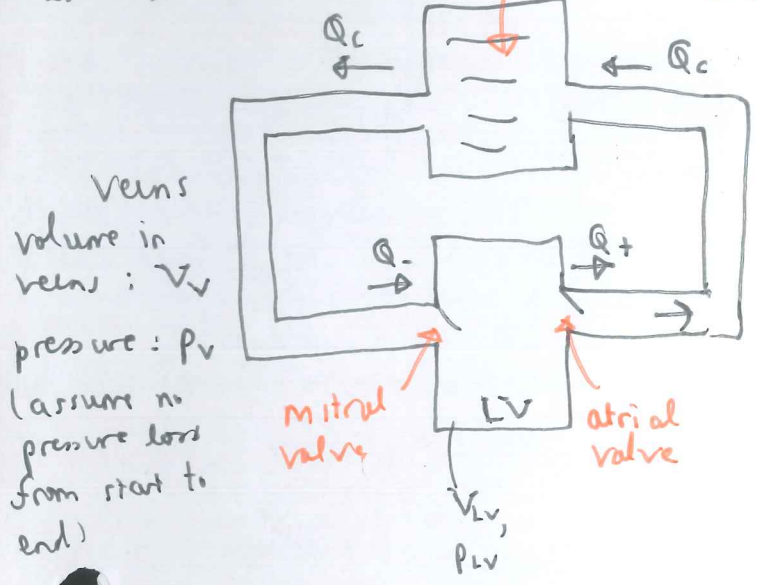
Low c = tight
High c = loose.



- A. The systole: Isovolumetric contraction. The compliance falls as the heart tightens.
- B. Ejection. Constant low compliance C_d (tight) pushes blood out.
- C. The diastole: Isovolumetric contraction. The compliance rises as the heart loosens.
- D. Refilling. Constant high compliance C_s (loose) allows blood in.

A simple mechanical model of the circulation.

We would like to write down a mathematical model to explain the $P_{LV} - V_{LV}$ plot.
 Capillaries - offer resistance but take up no volume.



arteries
 volume in arteries: V_a
 pressure: p_a

$Q_+, Q_-, Q_c =$ blood flows
 $V_a, V_V, V_{LV} =$ compartment volumes

Veins
 volume in veins: V_V
 pressure: p_V
 (assume no pressure loss from start to end)

Blood is incompressible but blood vessels are compliant (so blood flows at different points along the network can differ from one another and change with time)

Conservation of blood:

$$\dot{V}_a = Q_+ - Q_-$$

$$\dot{V}_V = Q_c - Q_-$$

$$\dot{V}_{LV} = Q_+ - Q_c$$

Resistances:

Capillaries occupy no volume but offer resistance:

$$Q_c = \frac{p_a - p_V}{R_c}$$

$$Q_+ = \frac{[p_{LV} - p_a]_+}{R_a}$$

the positive part

$$Q_- = \frac{[p_V - p_{LV}]_+}{R_V}$$

Compliance.

Increasing pressure distends blood vessels:

$$V_a = V_{a0} + C_a p_a$$

$$V_V = V_{V0} + C_V p_V$$

$$V_{LV} = V_{LV0} + C_{LV} p_{LV}$$

$C =$ compliance
 $= \frac{1}{\text{elastance}}$

low $c =$ tight, high $c =$ loose

Writing the model in terms of just pressures:

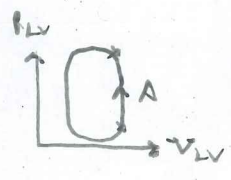
$$\frac{dp_a}{dt} = \frac{[P_{LV} - p_a]_+}{R_a C_a} - \frac{p_a - p_v}{R_c C_a} \quad (1)$$

$$\frac{dp_v}{dt} = \frac{p_a - p_v}{R_c C_v} - \frac{[p_v - P_{LV}]_+}{R_v C_v} \quad (2)$$

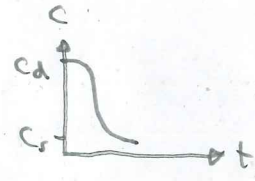
$$\frac{d(C_{LV} P_{LV})}{dt} = \frac{[p_v - P_{LV}]_+}{R_v} - \frac{[P_{LV} - p_a]_+}{R_a} \quad (3)$$

A. Isovolumetric contraction

Initial conditions:
 $P_{LV} < p_v \ll p_a$



(Both valves closed) (0.05s)



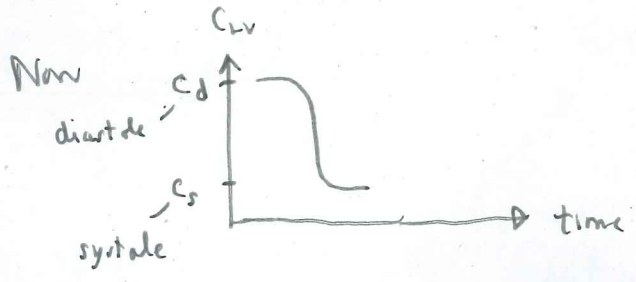
① ⇒ $\frac{dp_a}{dt} = - \frac{p_a}{R_c C_a}$

● $R_c C_a \approx 1.8s$ and time of contraction $\approx 0.05s$ so $p_a \approx \text{constant}$

on this phase

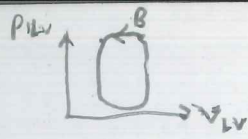
similarly, ② ⇒ $p_v \approx \text{constant}$ on this phase

③ ⇒ $\frac{d(C_{LV} P_{LV})}{dt} = 0$



so C_{LV} falls, P_{LV} rises

B. Ejection



(Atrial valve open) (0.3 s) ($C_{LV} = \text{constant} = C_s$) 77

$P_V \ll P_a < P_{LV}$ (because now P_{LV} is much higher)

① \Rightarrow

$$\frac{dP_a}{dt} = \frac{P_{LV} - P_a}{R_a C_a} - \frac{P_a - P_V}{R_e C_e} \quad (*)$$

$0.3s$ $0.09s$ $1.8s$

much smaller than the other two

\Rightarrow $P_{LV} = P_a$

② \Rightarrow

$$\frac{dP_V}{dt} = \frac{P_a - P_V}{R_e C_e} \Rightarrow$$

$P_V = \text{constant}$

$0.3s$ $60s$

③ \Rightarrow

$$\frac{d(C_{LV} P_{LV})}{dt} = - \frac{P_{LV} - P_a}{R_a}$$

Now $C_{LV} = \text{constant} = C_s$ substituting this into (*) gives

$$\frac{dP_a}{dt} = - \frac{C_s}{C_a} \frac{dP_a}{dt} - \frac{P_a - P_V}{R_e C_a}$$

$$(C_a + C_s) \frac{dP_a}{dt} = - \frac{P_a}{R_e} \quad (\text{since } P_a \gg P_V)$$

$P_a \propto \exp \left[\frac{-t}{R_e (C_a + C_s)} \right]$

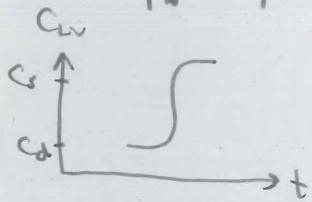
$0.2s$

$2.2s$

\Rightarrow P_a falls by ~ 0.87 in this phase.

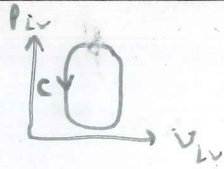
C. Isovolumetric relaxation

$$P_v < P_{LV} < P_a$$

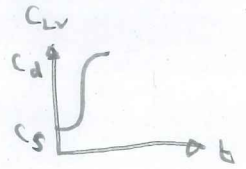


$$P_a = \text{constant}$$

$$P_v = \text{constant}$$



(Both valves closed) (0.08s)



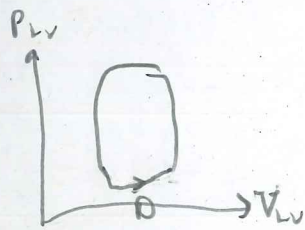
\$C_{LV}\$ rises.

As before, $\frac{d(C_{LV} P_{LV})}{dt} = 0$

So now P_{LV} falls until $P_{LV} = P_v$ and mitral valve open to commence filling

D. Refilling

$$P_{LV} < P_v < P_a$$



(Mitral valve open) (0.5s) ($C_{LV} = \text{constant} = C_d$)

① $\Rightarrow \frac{dP_a}{dt} = - \frac{P_a - P_v}{R_c C_a} = - \frac{P_a}{R_c C_a}$ since $P_a \gg P_v$

② $\Rightarrow \frac{dP_v}{dt} = \frac{P_a - P_v}{R_c C_v} - \frac{P_v - P_{LV}}{R_v C_v}$

③ $\Rightarrow \frac{d(P_v - P_{LV})}{dt} = \frac{P_v - P_{LV}}{C_d R_v}$ (since $C_v = \text{constant} = C_d$)

④ $\Rightarrow P_a \propto \exp\left[-\frac{t}{R_c C_a}\right] \Rightarrow P_a$ falls by ~ 0.76 in this phase

So total fall in P_a is $0.87 > 0.76 = 0.11$ (ejection) (refilling)

(120 mm Hg to 80 mm Hg)

① + ② gives

$$\dot{p}_v - \dot{p}_{Lv} = - \left(\frac{1}{R_v C_v} + \frac{1}{R_v C_A} \right) (p_v - p_{Lv})$$

since
($C_v = \text{constant}$)

$$p_v - p_{Lv} \propto \exp \left[- \frac{1}{R_v} \left(\frac{1}{C_v} + \frac{1}{C_A} \right) t \right]$$

Nervous control of the heart

This is what controls the heart rate, stroke volume and arterial blood pressure.

There are two parts of the nervous system that control cardiac output - the sympathetic and parasympathetic systems.

Nervous control is effected by - afferent nerves (to the brain)

efferent nerves (from the brain)

The sympathetic system releases noradrenaline and adrenaline and other neurotransmitters

There are two parts to the sympathetic system -

- α -sympathetic (peripheral vessels)
- β -sympathetic (ventricular muscle)

release of neurotransmitters here causes vasoconstriction which causes an increase in blood pressure.

release of neurotransmitters here causes an increase in the firing of the SA node
(ie increase in heart rate)

The sympathetic system acts slowly (~10s)

The parasympathetic system releases acetylcholine (another neurotransmitter)

This decrease the heart rate and causes vasodilation

This acts quickly.

The baroreceptors control the blood flow and blood pressure.

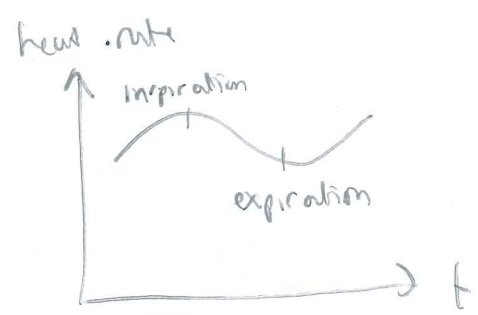
They are located in the aortic arch of the chest and the carotid sinus in the chest. The control response is called the baro reflex.

Oscillatory patterns

Respiratory sinus arrhythmia (RSA)

- heart rate is faster breathing in than breathing out
 (can get a different resting heart rate depending on which of these you are doing while measuring it)

reason:
 - inspiration leads to low pressure which increases the heart rate because it is easier to flow



Mayer waves

- due to the sympathetic system.
 - ~10s time period (same as timescale of sympathetic system).

Mathematical model of the baroreflex: the Otterén model (1997)

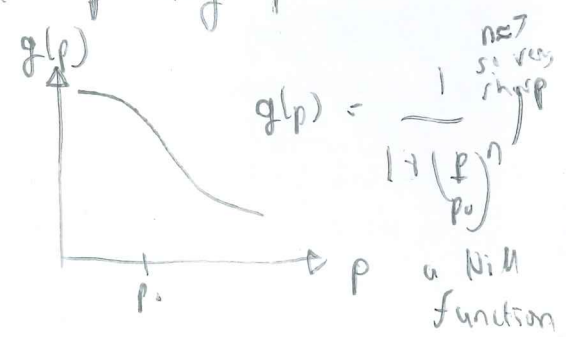
Three variables: average arterial and venous pressures p_a and p_v and the heart rate

Control is effected by sympathetic and parasympathetic tones:

Sympathetic tone:
 $T_s = g(p_a^r)$

$p_a^r = p_a(t - \tau)$

(a delay because its effects are delayed (after release - slow))



Parasympathetic tone : $T_p = 1 - g(p_a)$

Heart rate, H : $\dot{H} = \delta_H (N_0 - H) + \lambda_N T_s - \mu_N T_p$

(taken as a continuous variable - i.e. the average heart rate)

Natural resting heart rate in the absence of tones

Parameters denoting the strength of the sympathetic and parasympathetic tones

$$C_a \dot{p}_a = - \frac{p_a}{R_c} + H \Delta V$$

Compliance (as introduced before)
resistance as introduced before
stroke volume

Non-dimensionalization

$N = N_0 h, \quad p_a = p_0 p, \quad t = \tau \hat{t}$

$\Rightarrow \dot{p} = k(-p + \nu h)$

$\varepsilon \dot{h} = \delta(1-h) + \lambda g(p_i) - \mu(1-g(p))$

$p_i = p(t-1), \quad g(p) = \frac{1}{1+p^n} \quad n \approx 7$

($\varepsilon \ll 1$)
(other parameters roughly 0.1)