# Null space and Linear Programming

Mark Poolman

January 17, 2018

Wednesday L5



Null space: Encapsulates all possible steady-state solutions.

Enzyme subsets: Sets of reactions carrying flux in fixed ratio.

Elementary modes: Minimal, independent pathways in a system



Null space: Encapsulates all possible steady-state solutions.

Enzyme subsets: Sets of reactions carrying flux in fixed ratio.

Elementary modes: Minimal, independent pathways in a system



Null space: Encapsulates all possible steady-state solutions.

Enzyme subsets: Sets of reactions carrying flux in fixed ratio.

Elementary modes: Minimal, independent pathways in a system



Null space: Encapsulates all possible steady-state solutions.

Enzyme subsets: Sets of reactions carrying flux in fixed ratio.

Elementary modes: Minimal, independent pathways in a system



Null space: Encapsulates all possible steady-state solutions.

Enzyme subsets: Sets of reactions carrying flux in fixed ratio.

Elementary modes: Minimal, independent pathways in a system



Null space: Readily calculated, but can't analyse by

inspection.

Enzyme subsets: Not as generally useful compared to small

models.

Elementary modes: Impractical.



Null space: Readily calculated, but can't analyse by inspection.

Enzyme subsets: Not as generally useful compared to small models

Elementary modes: Impractical.



Null space: Readily calculated, but can't analyse by inspection.

Enzyme subsets: Not as generally useful compared to small models.

Elementary modes: Impractical.



Null space: Readily calculated, but can't analyse by inspection.

Enzyme subsets: Not as generally useful compared to small models.

Elementary modes: Impractical.



Null space: Readily calculated, but can't analyse by inspection.

Enzyme subsets: Not as generally useful compared to small models.

Elementary modes: Impractical.



# Other disadvantages of null-space analysis

- Provides a rather 'unfocussed' view of the system.
- Does not (implicitly) take into account thermodynamics.
- Hard to integrate experimental flux observations.
- Less interpretable for large (genome-scale) models.
- (Still very useful for validation).

Linear programming calculates a specific solution to the equation:

$$Nv=0\\$$

Subject to some additional information supplied by the user - at least one flux value specified.

```
\begin{array}{lll} \text{min/max} & : & \textbf{V}_{\text{targs}} & \longleftarrow \text{ objective} \\ \text{subject to} & \begin{cases} \textbf{N}\textbf{v} = \textbf{0} & \longleftarrow \text{ steady state} \\ \text{max}_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{ flux constraints} \end{cases}
```

- Maximise output(s) (need to fix input(s))
- FBA maximise growth rate for fixed input.
- Minimise input(s) (need to fix output(s))
- Minimise all reactions (need to fix input(s) and/or output(s))



$$\begin{array}{lll} \min/\max & : \quad \textbf{V}_{targs} & \longleftarrow \text{ objective} \\ \text{subject to} & \begin{cases} \textbf{N}\textbf{v} = \textbf{0} & \longleftarrow \text{ steady state} \\ \max_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{ flux constraints} \end{cases}$$

- Maximise output(s) (need to fix input(s))
- FBA maximise growth rate for fixed input.
- Minimise input(s) (need to fix output(s))
- Minimise all reactions (need to fix input(s) and/or output(s))



$$\begin{array}{lll} \min/\max & : \quad \textbf{V}_{\text{targs}} & \longleftarrow \text{ objective} \\ \text{subject to} & \begin{cases} \textbf{N}\textbf{v} = \textbf{0} & \longleftarrow \text{ steady state} \\ \max_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{ flux constraints} \end{cases}$$

- Maximise output(s) (need to fix input(s))
- FBA maximise growth rate for fixed input.
- Minimise input(s) (need to fix output(s))
- Minimise all reactions (need to fix input(s) and/or output(s))



$$\begin{array}{lll} \text{min/max} & : & \textbf{v}_{\text{targs}} & \longleftarrow \text{objective} \\ \text{subject to} & \begin{cases} \textbf{Nv} = \textbf{0} & \longleftarrow \text{steady state} \\ \text{max}_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{flux constraints} \end{cases}$$

- Maximise output(s) (need to fix input(s))
- FBA maximise growth rate for fixed input.
- Minimise input(s) (need to fix output(s))
- Minimise all reactions (need to fix input(s) and/or output(s))



```
\begin{array}{lll} \text{min/max} & : & \textbf{v}_{targs} & \longleftarrow \text{objective} \\ \text{subject to} & \left\{ \begin{array}{lll} \textbf{Nv} = \textbf{0} & \longleftarrow \text{steady state} \\ \text{max}_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{flux constraints} \end{array} \right. \end{array}
```

- Maximise output(s) (need to fix input(s))
- FBA maximise growth rate for fixed input.
- Minimise input(s) (need to fix output(s))
- Minimise all reactions (need to fix input(s) and/or output(s))



$$\begin{array}{lll} \text{min/max} & : & \textbf{v}_{targs} & \longleftarrow \text{objective} \\ \text{subject to} & \left\{ \begin{array}{ll} \textbf{N}\textbf{v} = \textbf{0} & \longleftarrow \text{steady state} \\ \text{max}_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{flux constraints} \end{array} \right. \end{array}$$

#### Typical flux constraints:

- $\min_{i} = \max_{i} \neq 0$ : flux is fixed
- $\min_i = \max_i = 0$ : reaction is knocked out.
- $\min_i = 0$ ,  $\max_i \neq 0$ : force irreversible L->R
- $\min_i \neq 0$ ,  $\max_i = 0$ : force irreversible R->L



## Advantages of FBA

Very fast.

Integrates flux data.

- Easy to reformulate the problem and solve again.
- The reactions in a solution can be extracted from the main model for more detailed analysis.

## Disadvantages of FBA

Only provides a single solution.

Potential for numerical instability (esp. if maximising).

Potential for multiple optima.

• Choice of the objective is subjective (!)

## Exploring the optimal space - constraint scanning

$$\begin{array}{lll} \text{minimise} & : & \textbf{v}_{targs} & \longleftarrow \text{objective} \\ \text{subject to} & \left\{ \begin{array}{ll} \textbf{N}\textbf{v} = \textbf{0} & \longleftarrow \text{steady state} \\ \text{max}_i \geq \textbf{v}_i \geq \min_i & \longleftarrow \text{flux constraints} \end{array} \right. \end{array}$$

- Find a solution.
- $\bullet$  Increment one (or more) of the constraints  $\boldsymbol{v}_i$
- Solve again.
- Repeat to build up a set of solutions.
- Identify correlated responses in the solution set.



## Example - identifying a catabolic core

A study of Salmonella spp.

Antibiotic challenges generate a stress response.

This increases the demand for ATP.

 How to identify which reactions will respond to this demand?

## Example - identifying a catabolic core

Scan over a range of ATP demand fluxes (while synthesising biomass) and identify responding reactions.

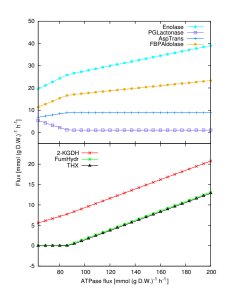
minimise : 
$$|\mathbf{v}|$$
  $\leftarrow$  objective - min. sum of fluxes   
 $\begin{cases} \mathbf{N}\mathbf{v} = \mathbf{0} & \leftarrow \text{ steady state constraint} \\ v_j = t_j & \leftarrow \text{ output transporters, constant} \\ v_{\text{ATPase}} = J_{\text{ATPase}} & \leftarrow \text{ ATP hydrolysis, varied} \end{cases}$ 

### Example - identifying a catabolic core

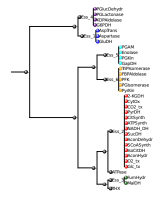
Scan over a range of ATP demand fluxes (while synthesising biomass) and identify responding reactions.

minimise : 
$$|\mathbf{v}|$$
  $\leftarrow$  objective — min. sum of fluxes   
subject to 
$$\begin{cases}
\mathbf{N}\mathbf{v} = \mathbf{0} & \leftarrow \text{ steady state constraint} \\
\mathbf{v}_j = t_j & \leftarrow \text{ output transporters, constant} \\
\mathbf{v}_{\text{ATPase}} = \mathbf{J}_{\text{ATPase}} & \leftarrow \text{ ATP hydrolysis, varied}
\end{cases}$$

# Results - flux response



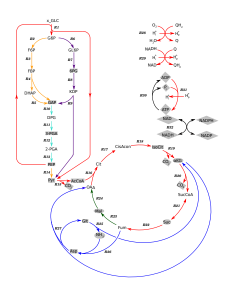
#### Results - flux correlations



33 reactions correlated with imposed ATPase.



#### Results - catabolic core



#### Results - condensed network

