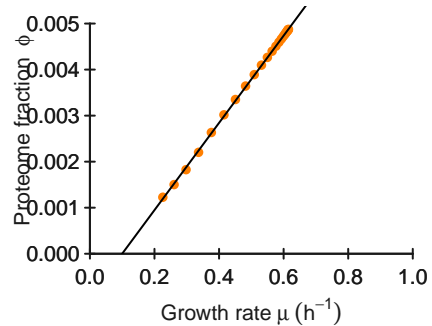
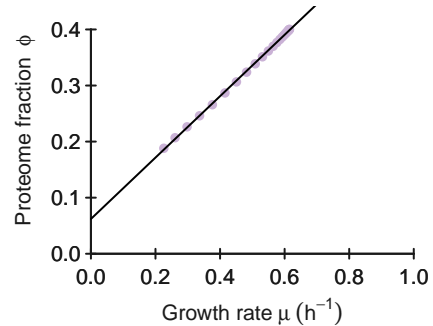
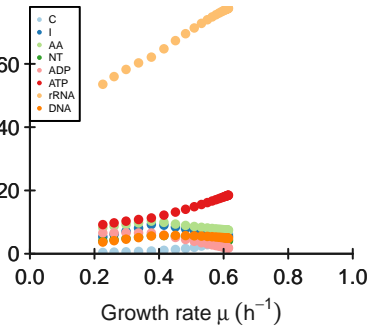
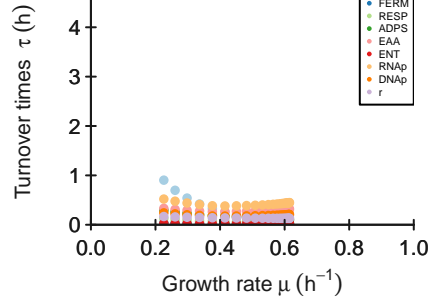
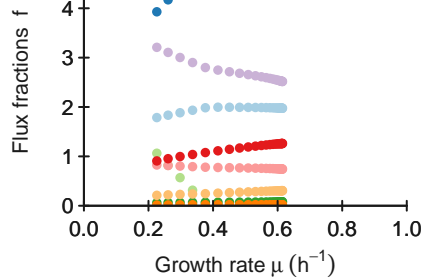
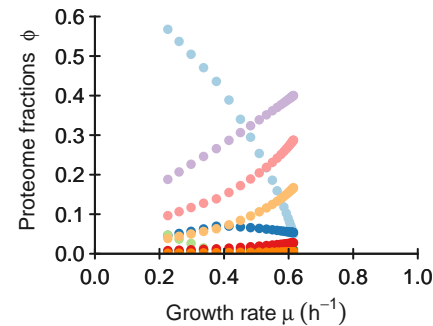
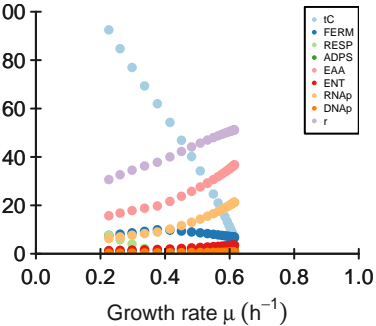
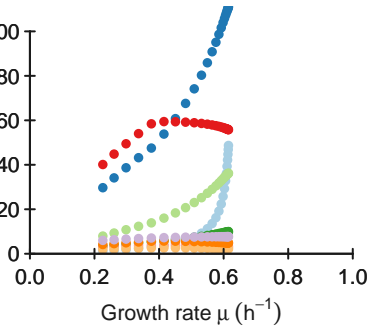
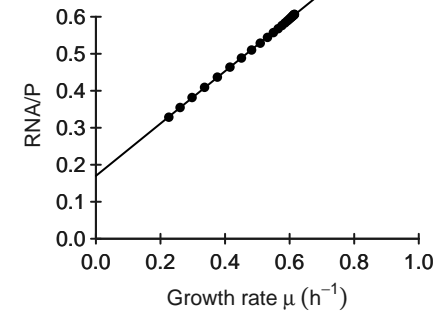
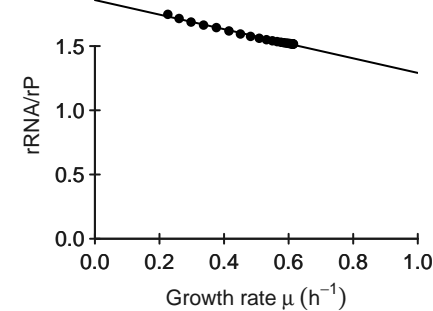
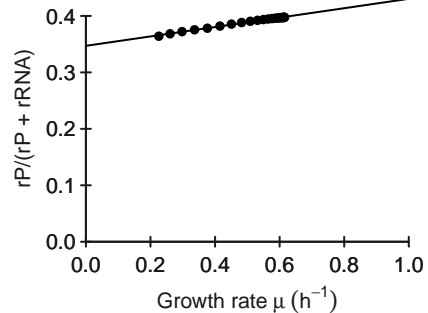
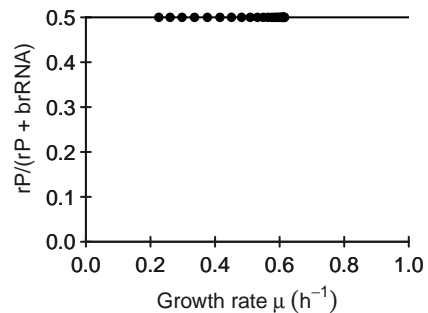


**DNAp****r**Metabolite concentrations  $c^m$  (g/L)Protein concentrations  $p$  (g/L)Apparent turnover numbers  $k_{app}$  ( $\text{h}^{-1}$ )**RNA/P****rRNA/rP**

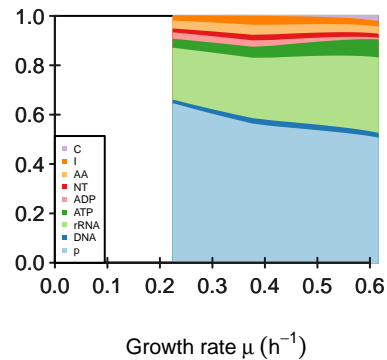
$rP/(rP + rRNA)$



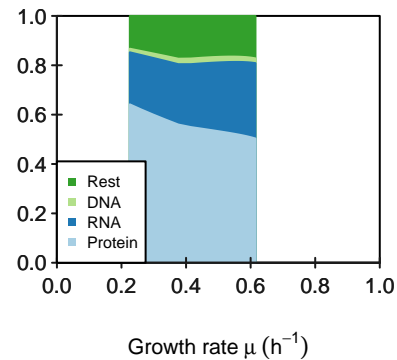
Protein mass fraction in ribosome



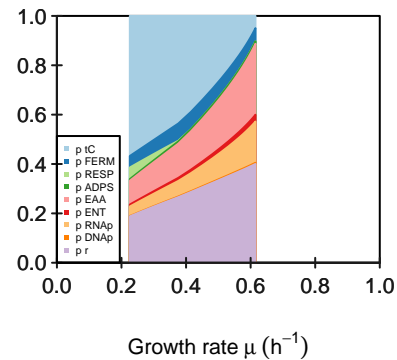
Relative biomass composition



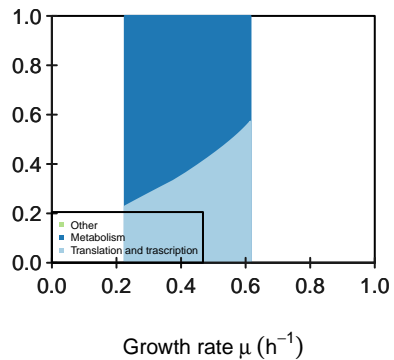
Predicted biomass



Proteome composition



Proteome sectors



keep\_ribosome\_kcat FALSE  
keep\_transport\_kcat FALSE  
maintenance\_fun constant

# M

[illegible]





# kcat

|       | tC | FERM  | RESP   | ADPS | EAA | ENT | RNAp | DNAP | r  |
|-------|----|-------|--------|------|-----|-----|------|------|----|
| kcatf | 56 | 373.5 | 186.75 | 26   | 7   | 149 | 6    | 13   | 19 |
| kcatb | 6  | 37    | 19     | 3    | 1   | 15  | 0    | 0    | 0  |



## Keq

| [1,] | [,1]              | [,2]             | [,3]             | [,4]             | [,5]             | [,6]             | [,7] | [,8] | [,9] |
|------|-------------------|------------------|------------------|------------------|------------------|------------------|------|------|------|
|      | 1306.666666666667 | 129.210810810811 | 125.810526315789 | 4.33333333333333 | 18.6666666666667 | 2.20740740740741 | Inf  | Inf  | Inf  |

## phi input

|      |       |       |       |       |       |       |       |       |      |
|------|-------|-------|-------|-------|-------|-------|-------|-------|------|
| [1,] | [,1]  | [,2]  | [,3]  | [,4]  | [,5]  | [,6]  | [,7]  | [,8]  | [,9] |
|      | 0.065 | 0.035 | 0.035 | 0.003 | 0.248 | 0.032 | 0.119 | 0.003 | 0.46 |

**average saturation input**

3

### minimal phi constraint

[illegible]

